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Hegy et al.

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(54) **LIGHT COLLECTION FROM OBJECTS WITHIN A FLUID COLUMN**

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G01N 15/10 (2006.01)

(52) **U.S. Cl.**
CPC **G01N 15/1459** (2013.01); **G01N 15/1436** (2013.01); **G01N 2015/1006** (2013.01)

(58) **Field of Classification Search**
CPC G01N 15/1459; G01N 15/1436; G01N 2015/1006
USPC 250/459.1
See application file for complete search history.

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Primary Examiner — David P Porta

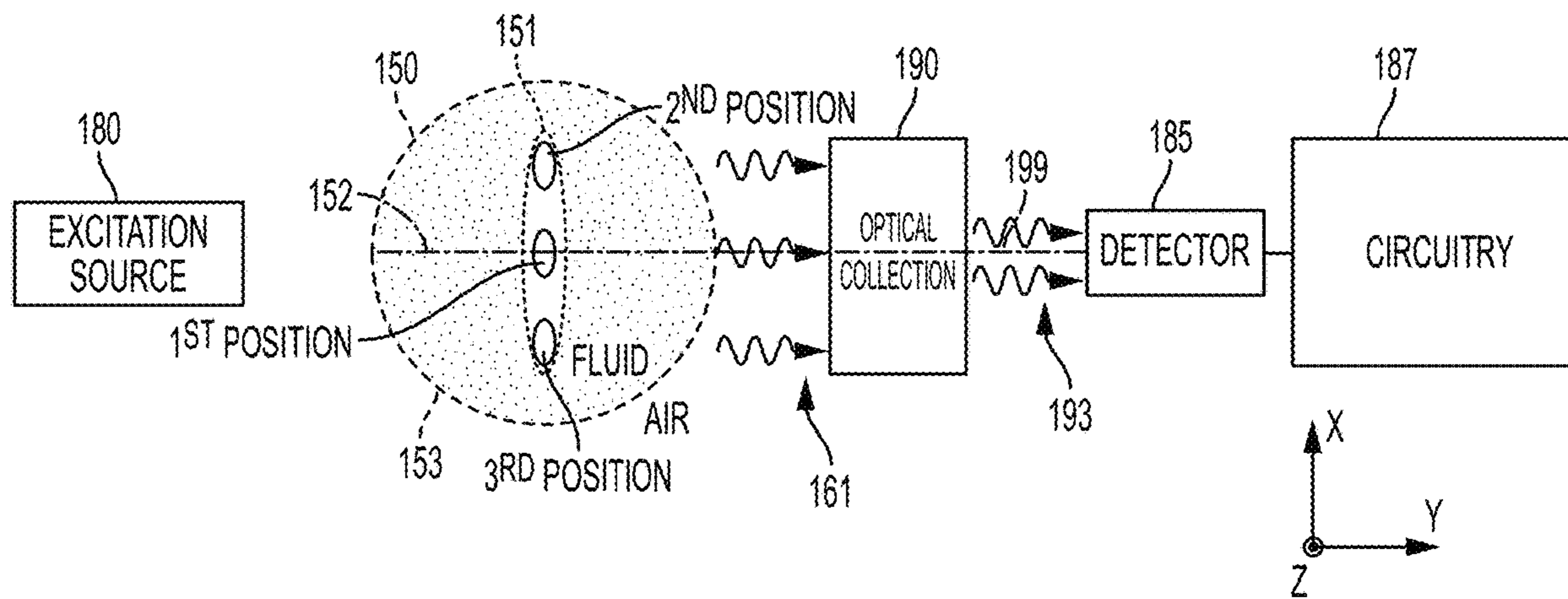
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(57) **ABSTRACT**

An optical arrangement receives output light emanating from an object disposed within a fluid column that crosses an optical refraction boundary of the fluid column between the object and the optical arrangement. The optical arrangement modifies the output light such that the modified output light has an intensity that is more uniform than the output light.

31 Claims, 25 Drawing Sheets



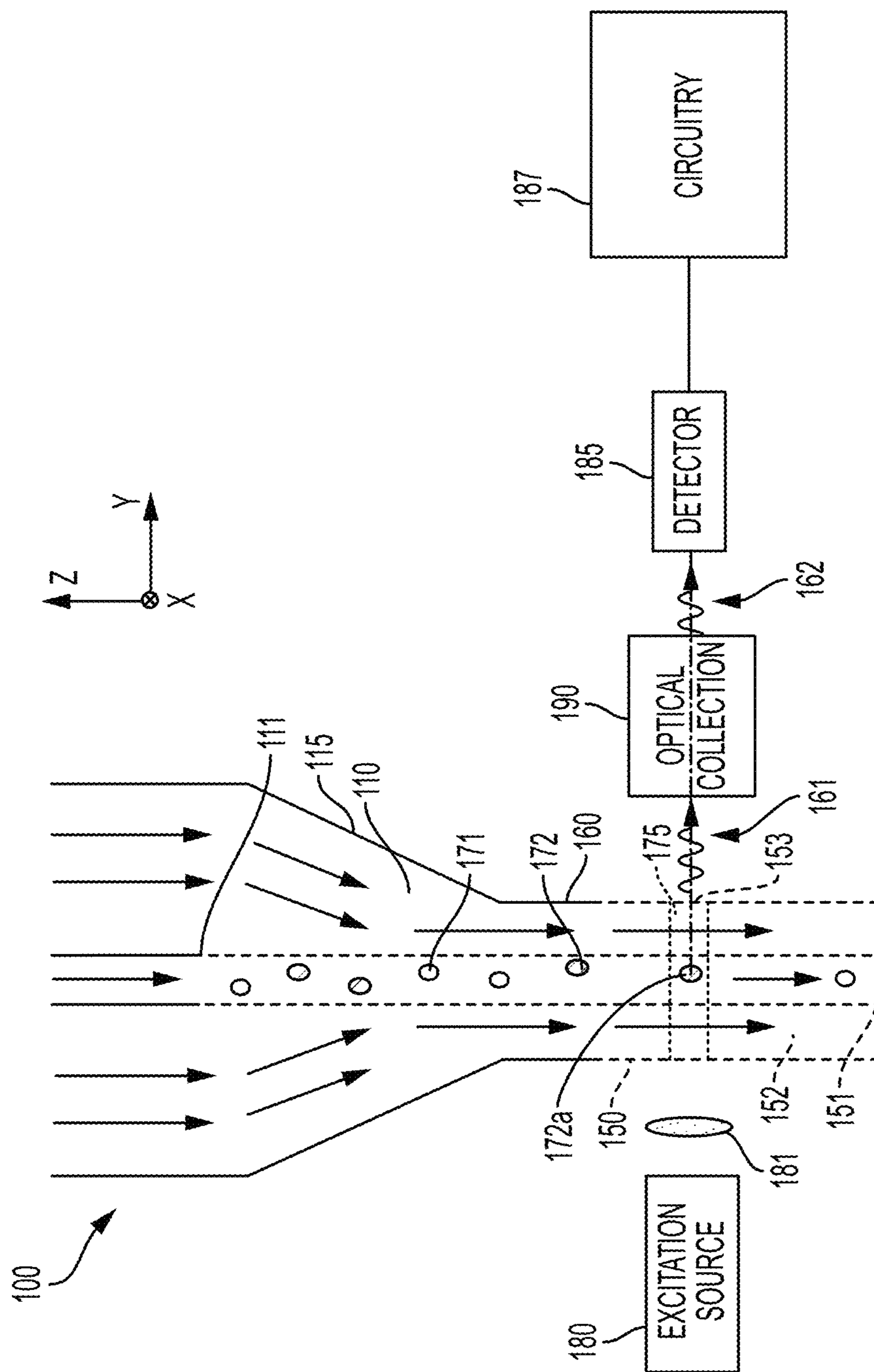


FIG. 1A

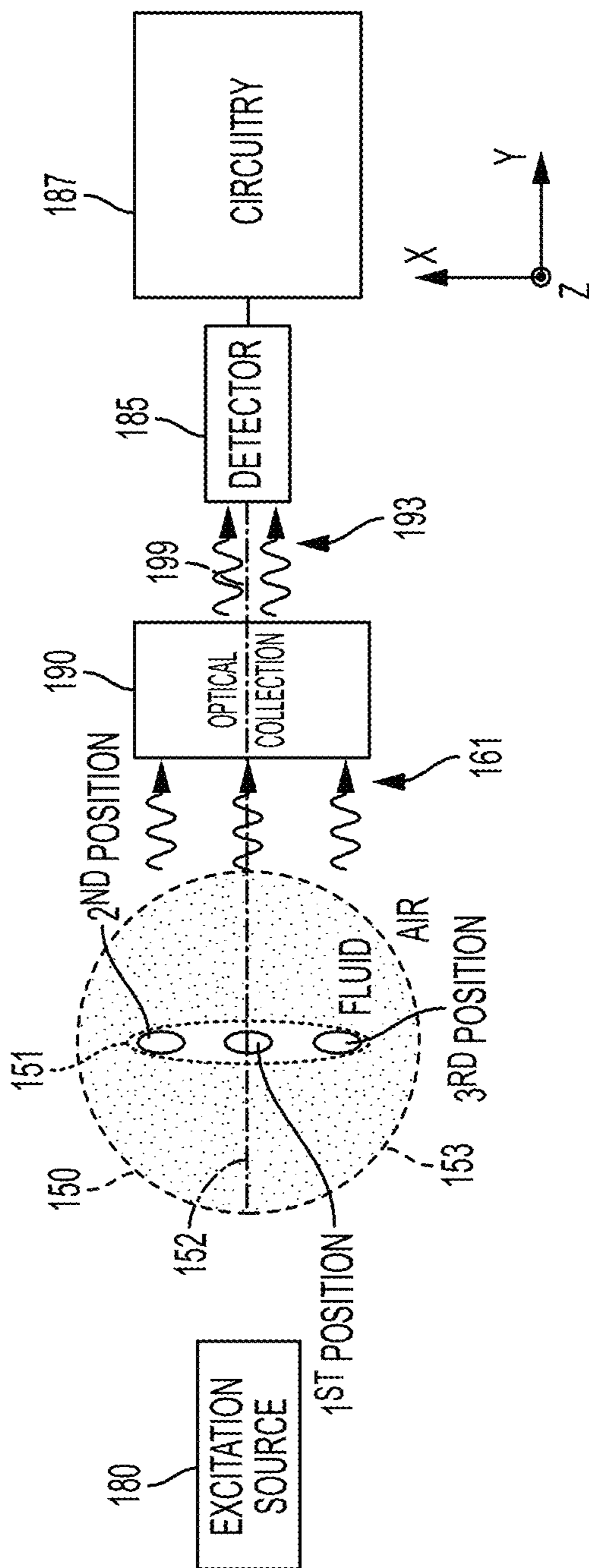


FIG. 1B

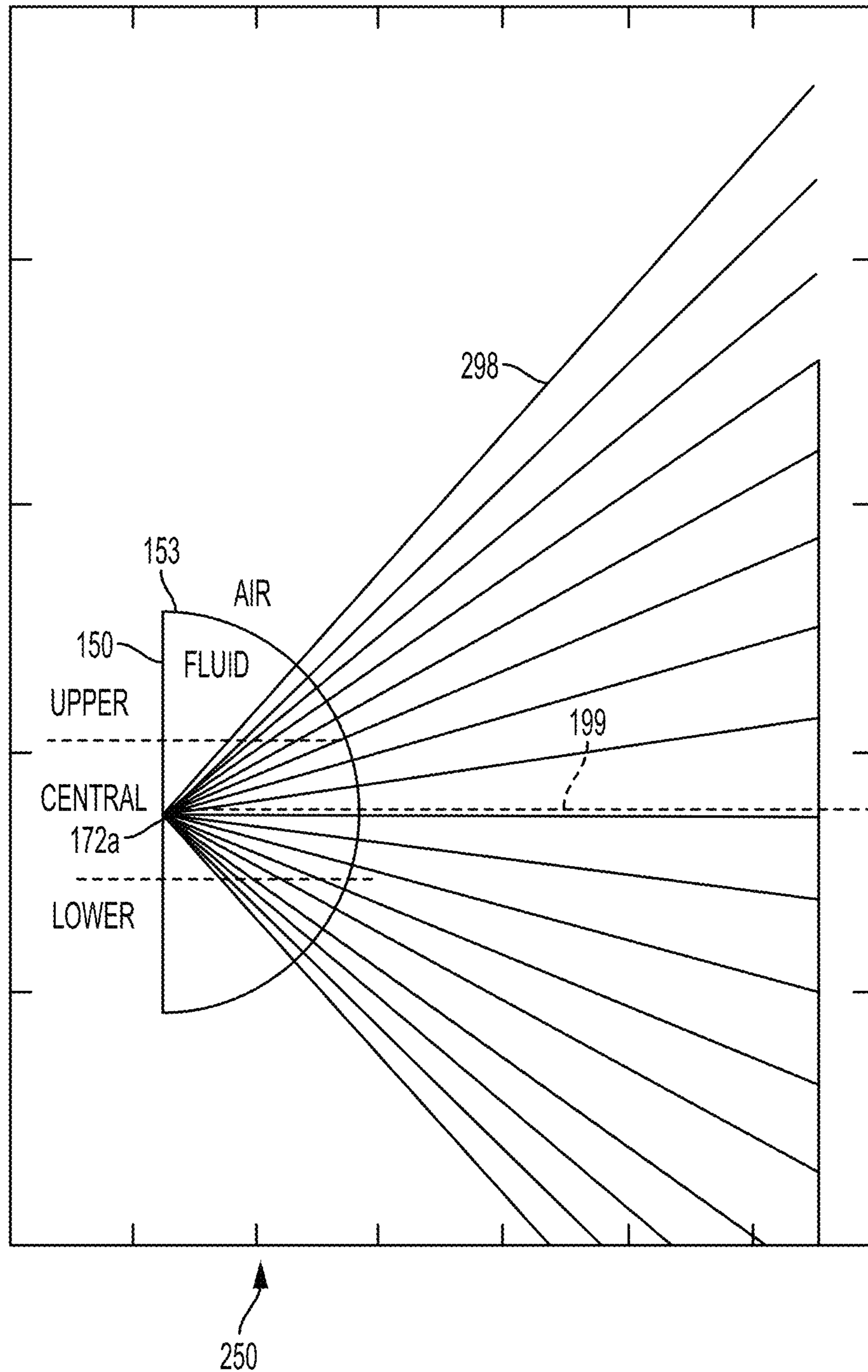


FIG. 2A

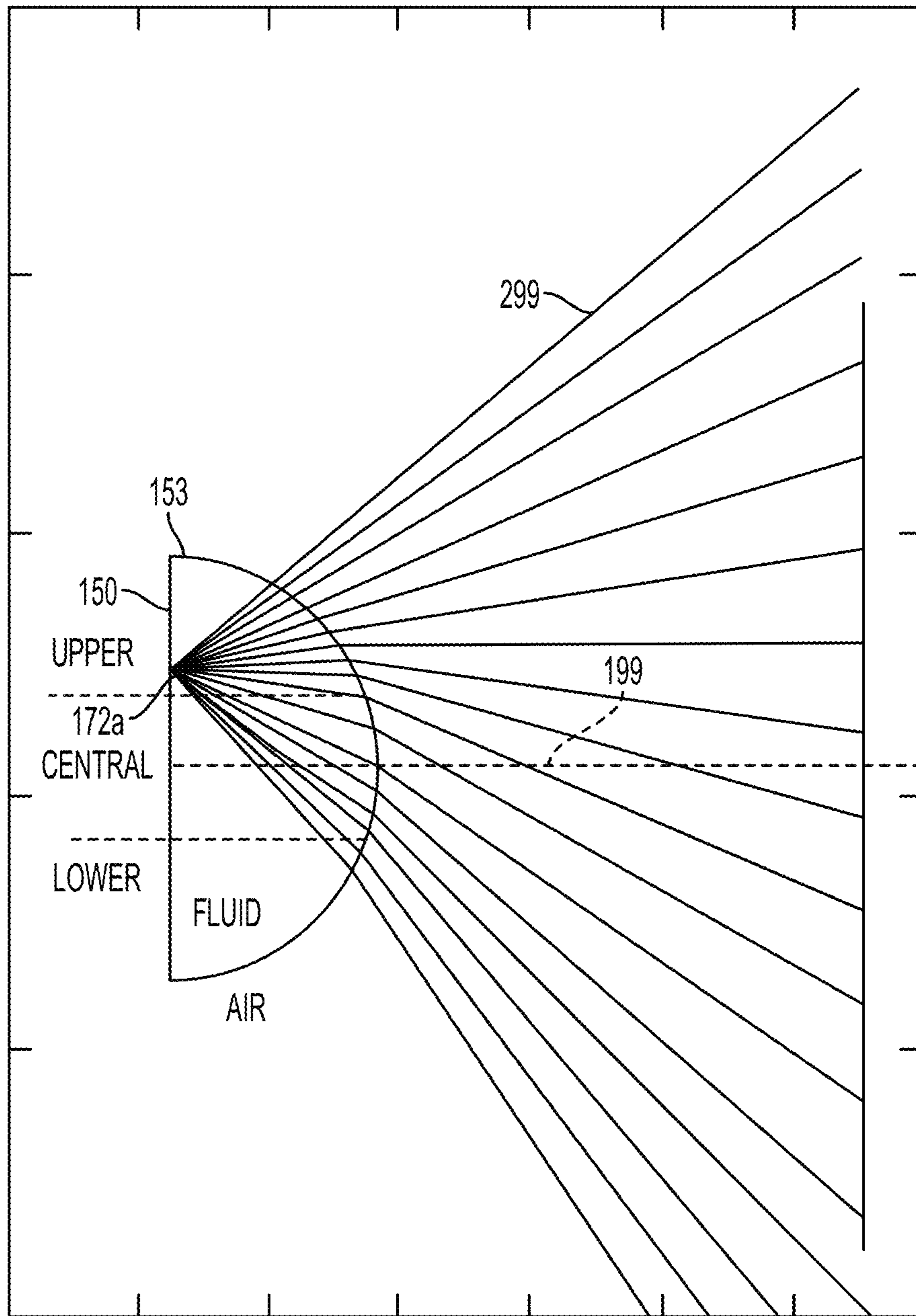


FIG. 2B

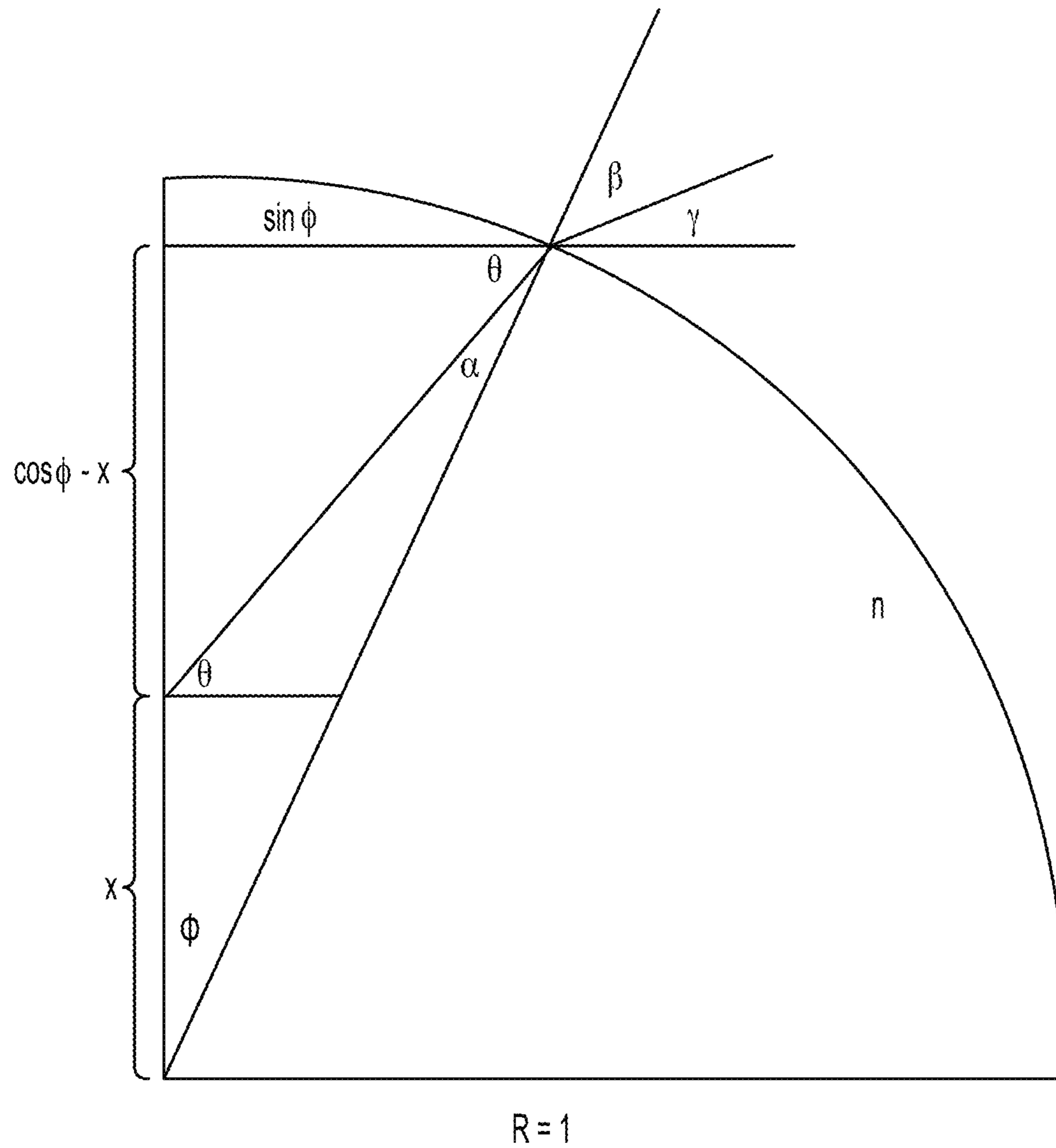


FIG. 3

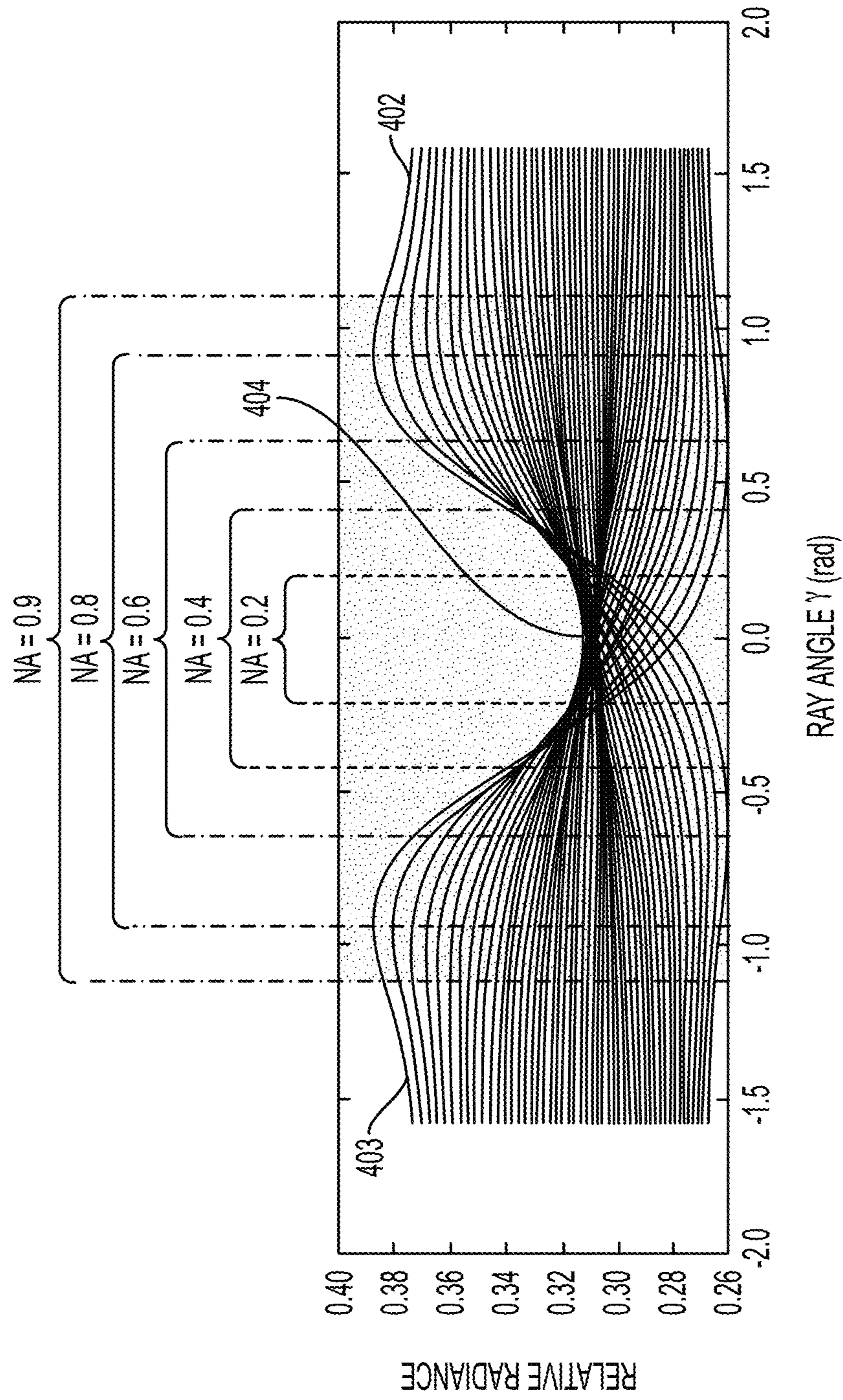


FIG. 4A

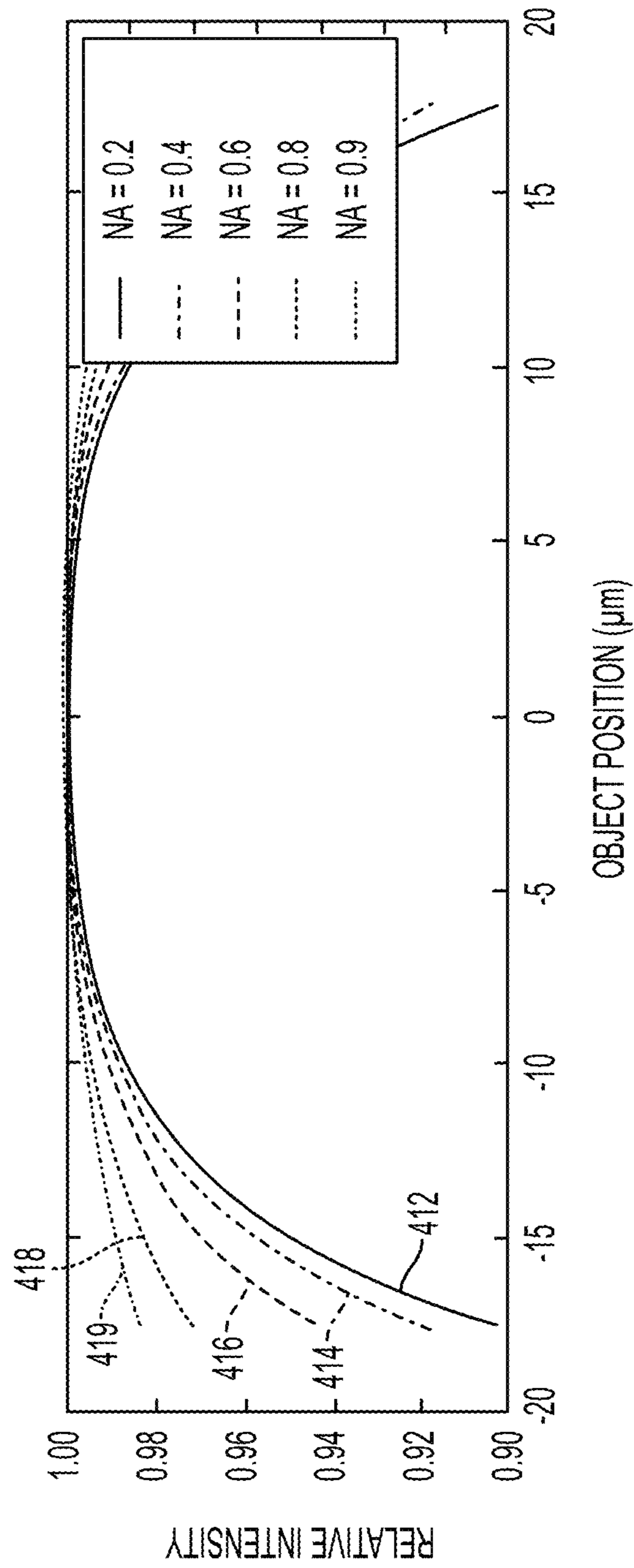


FIG. 4B

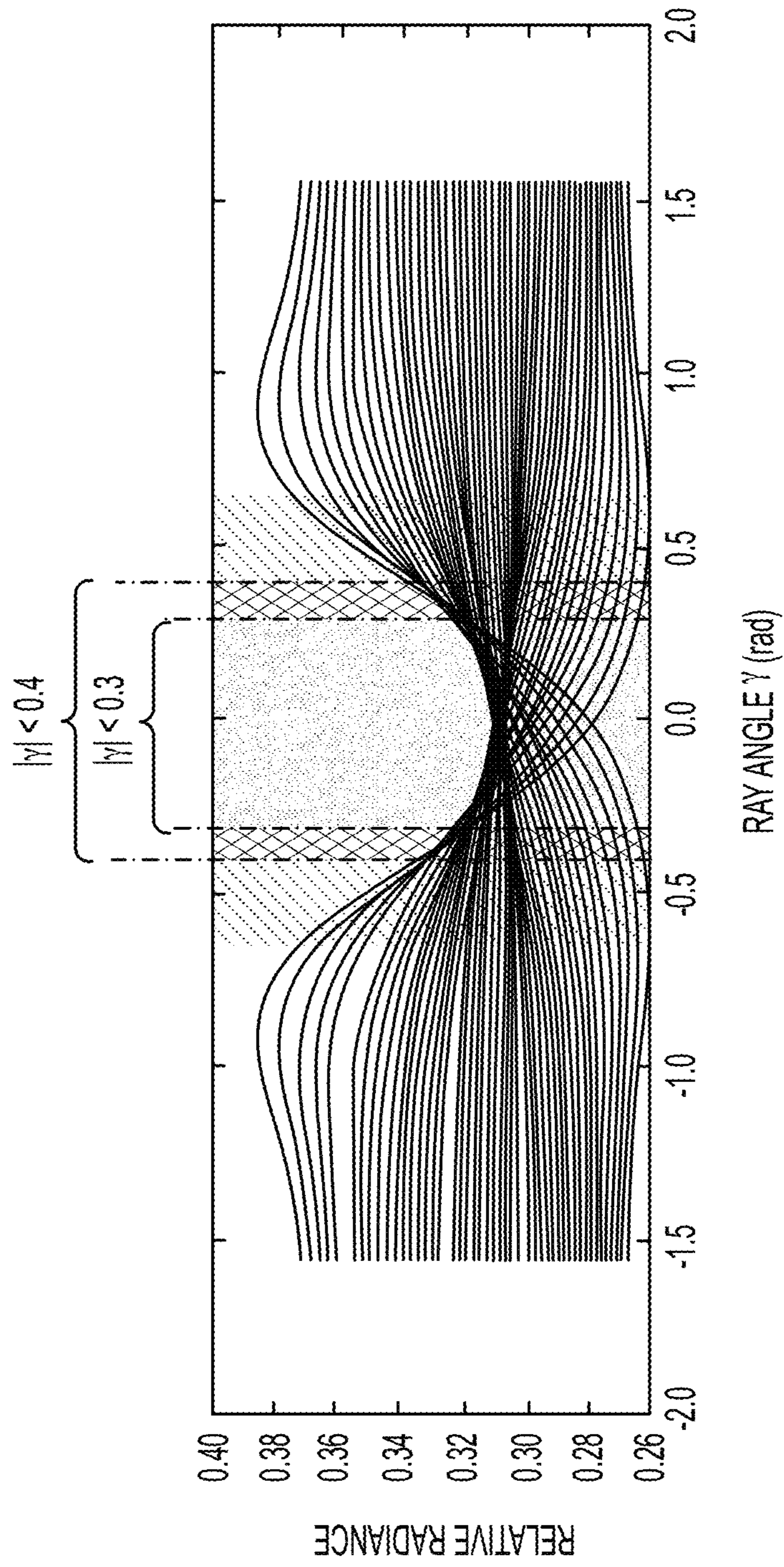


FIG. 5A

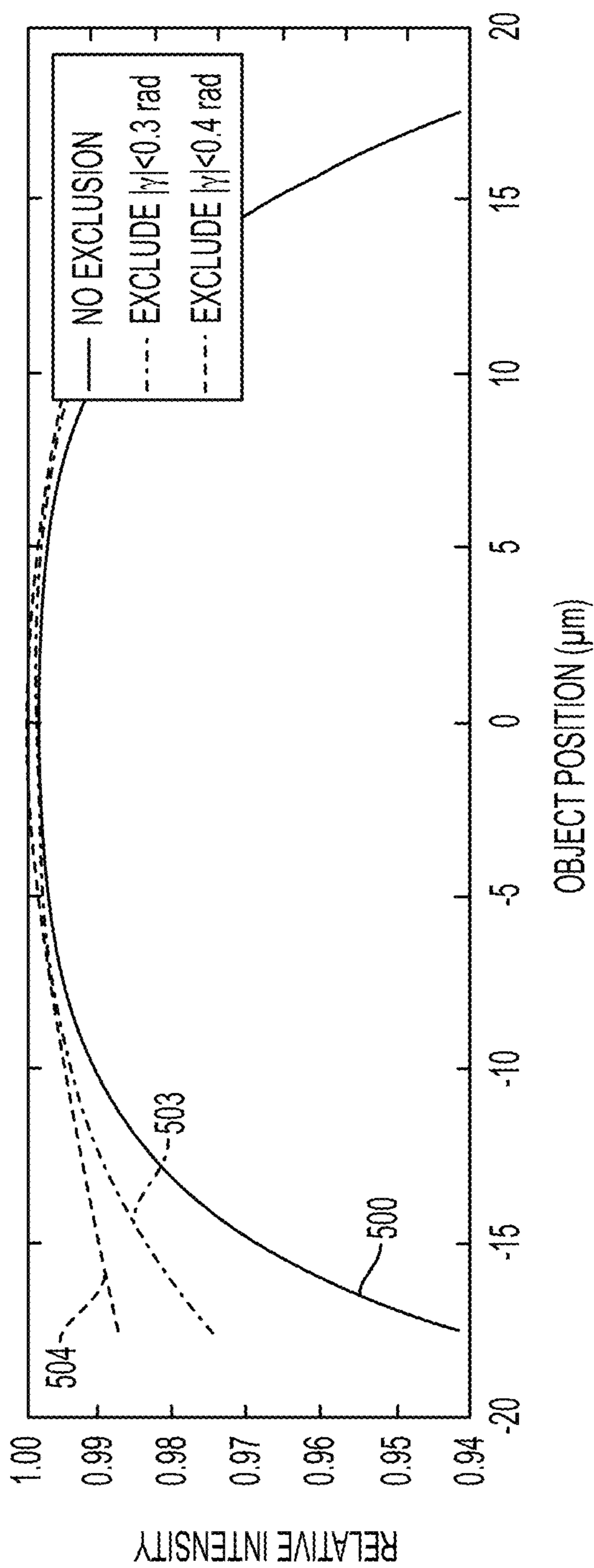


FIG. 5B

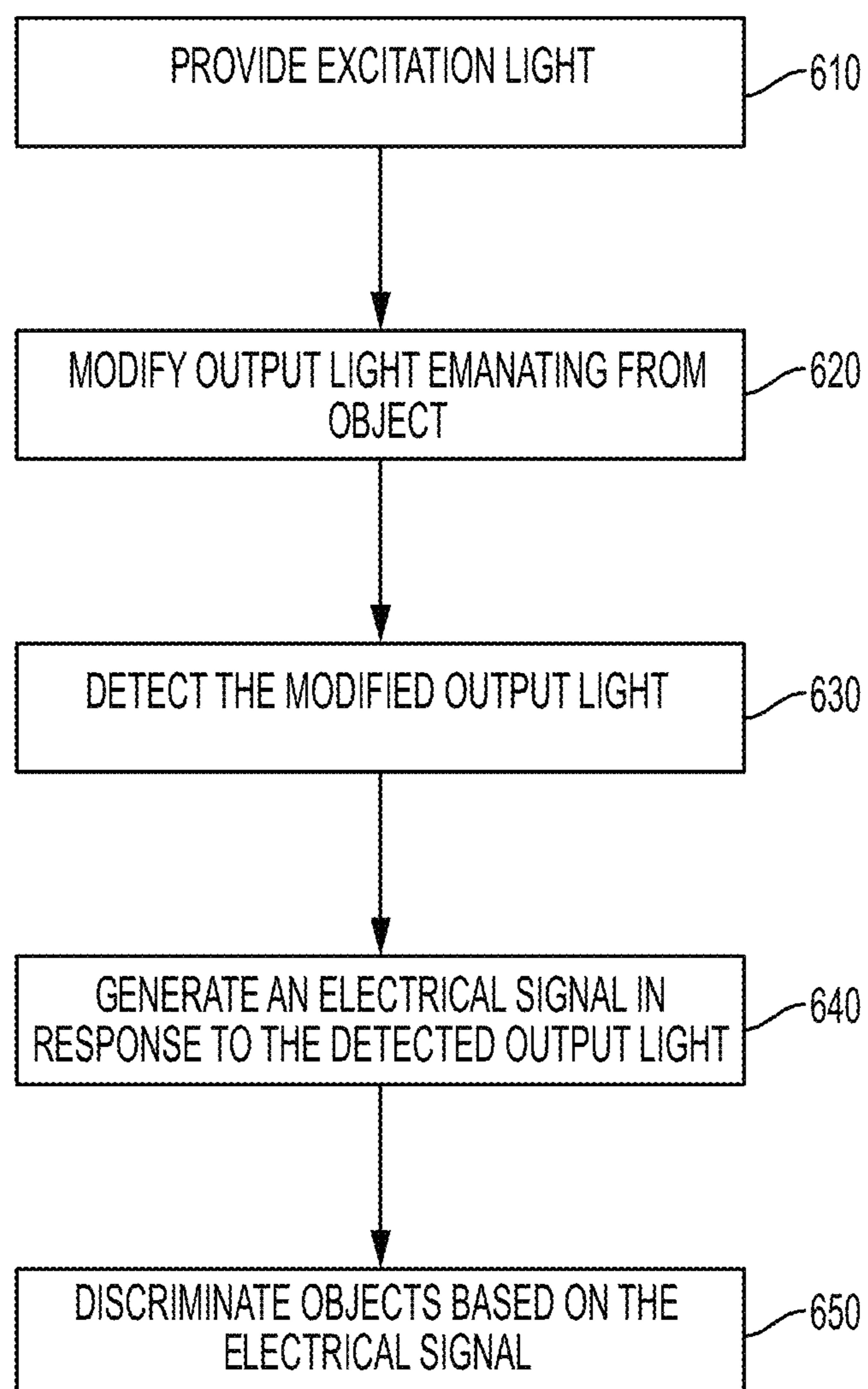


FIG. 6

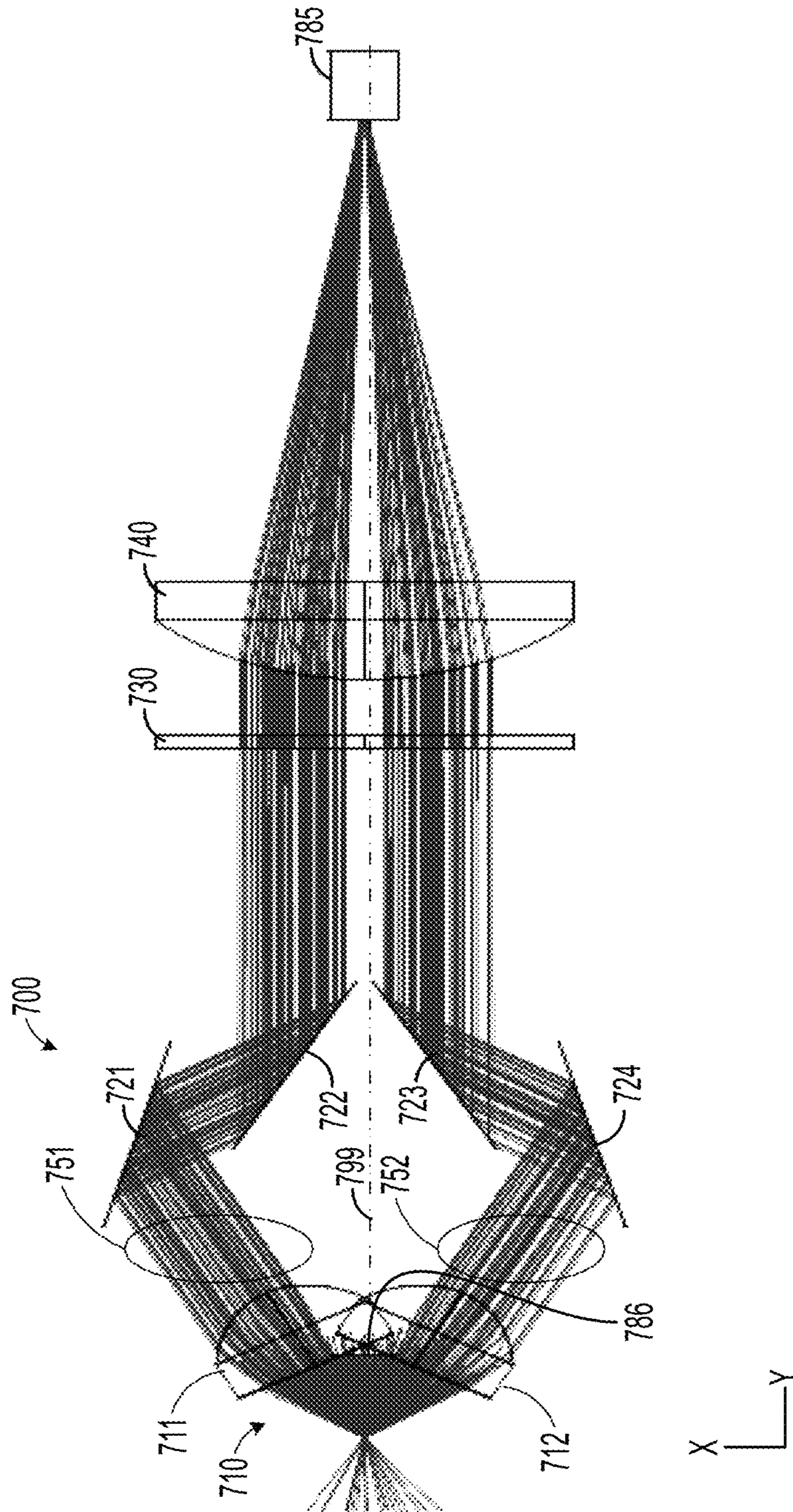


FIG. 7

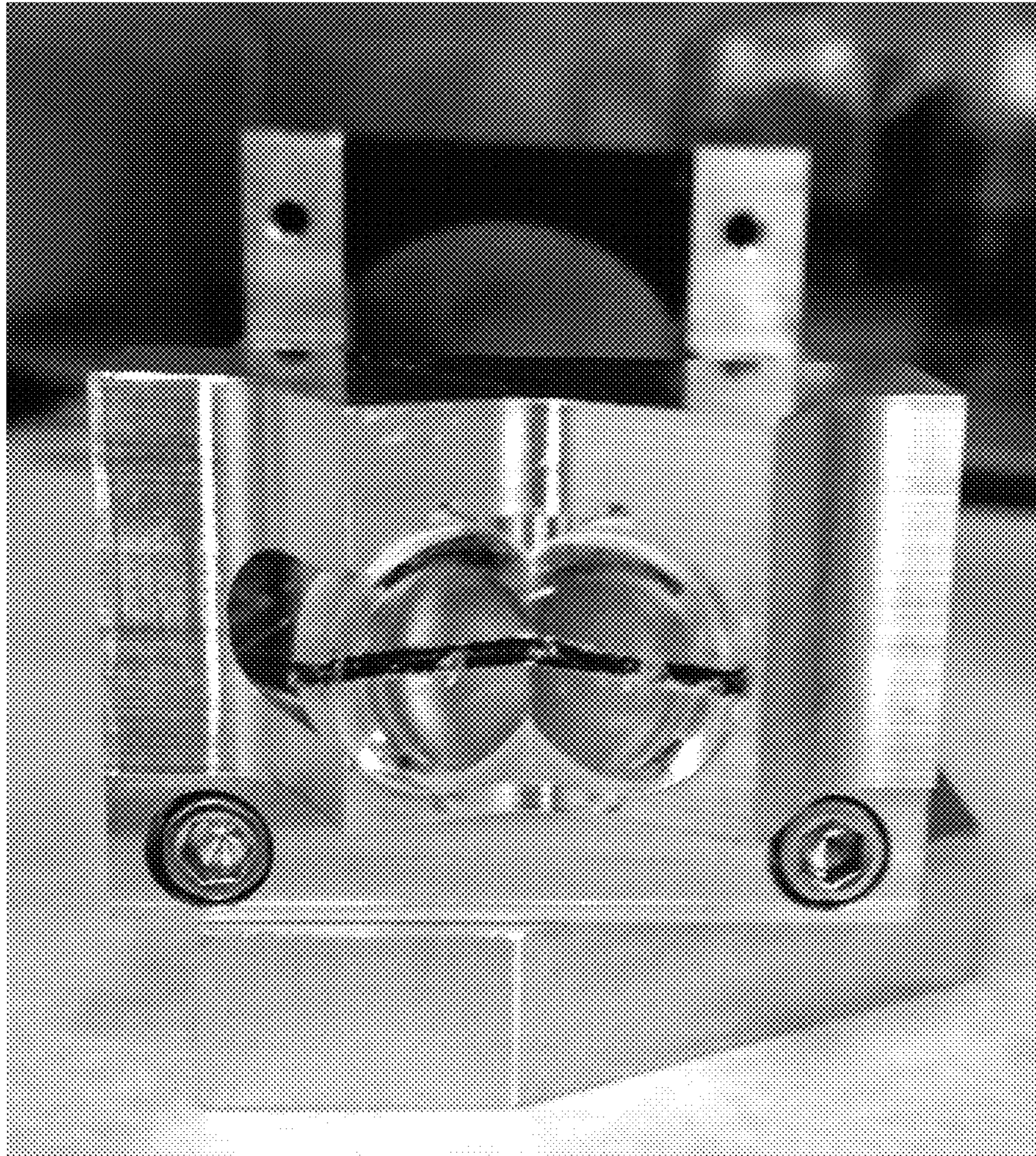


FIG. 8

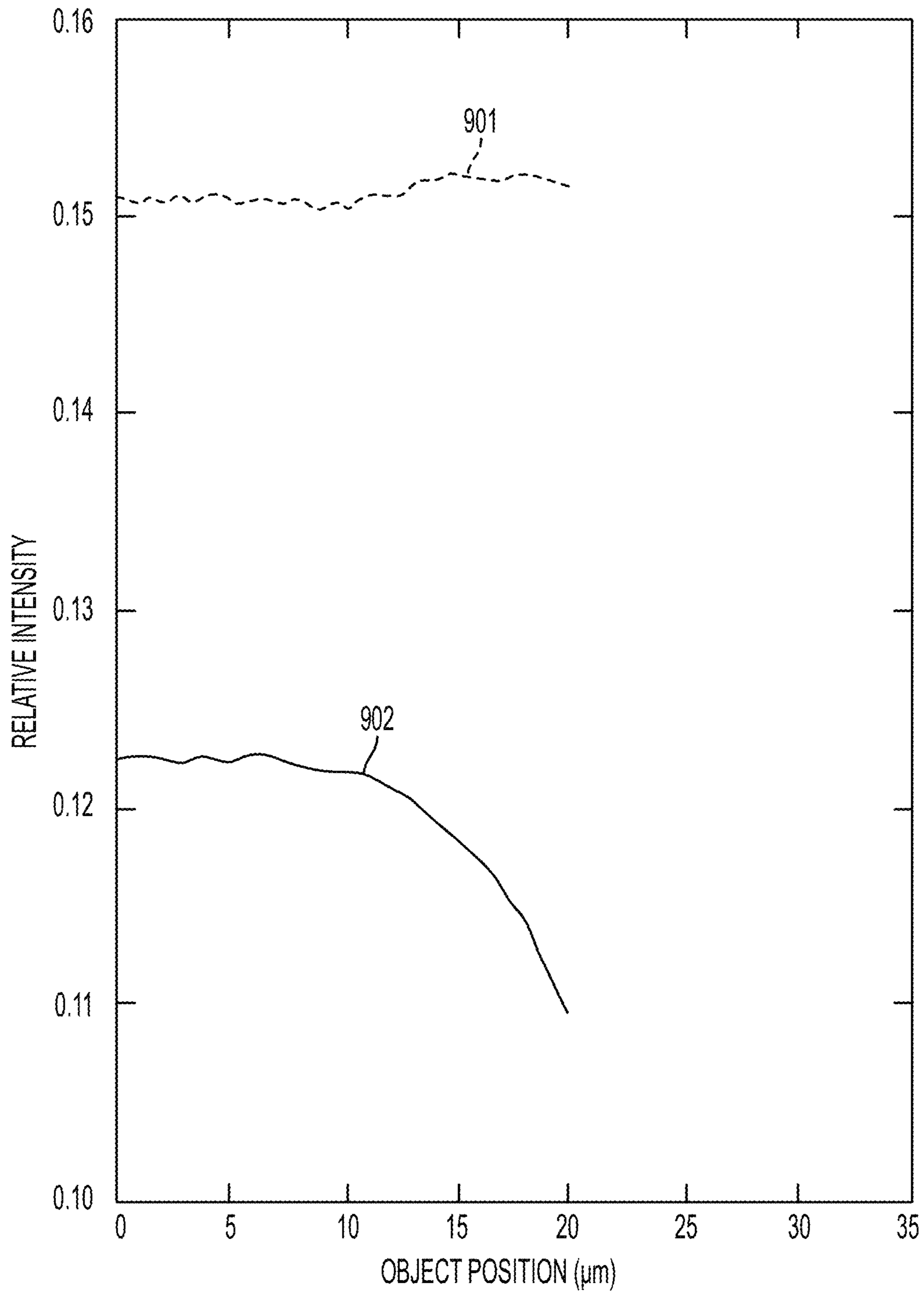


FIG. 9A

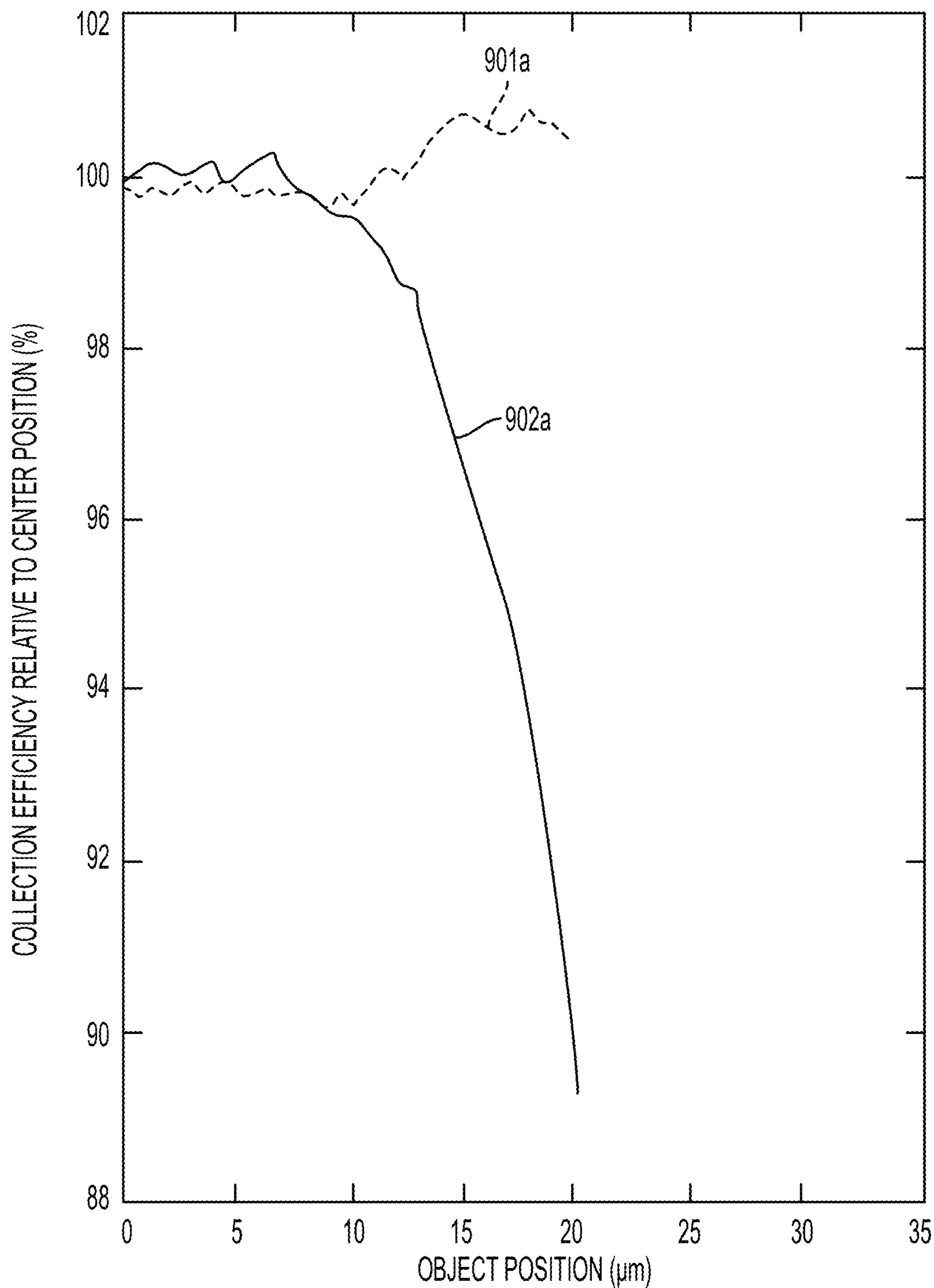


FIG. 9B

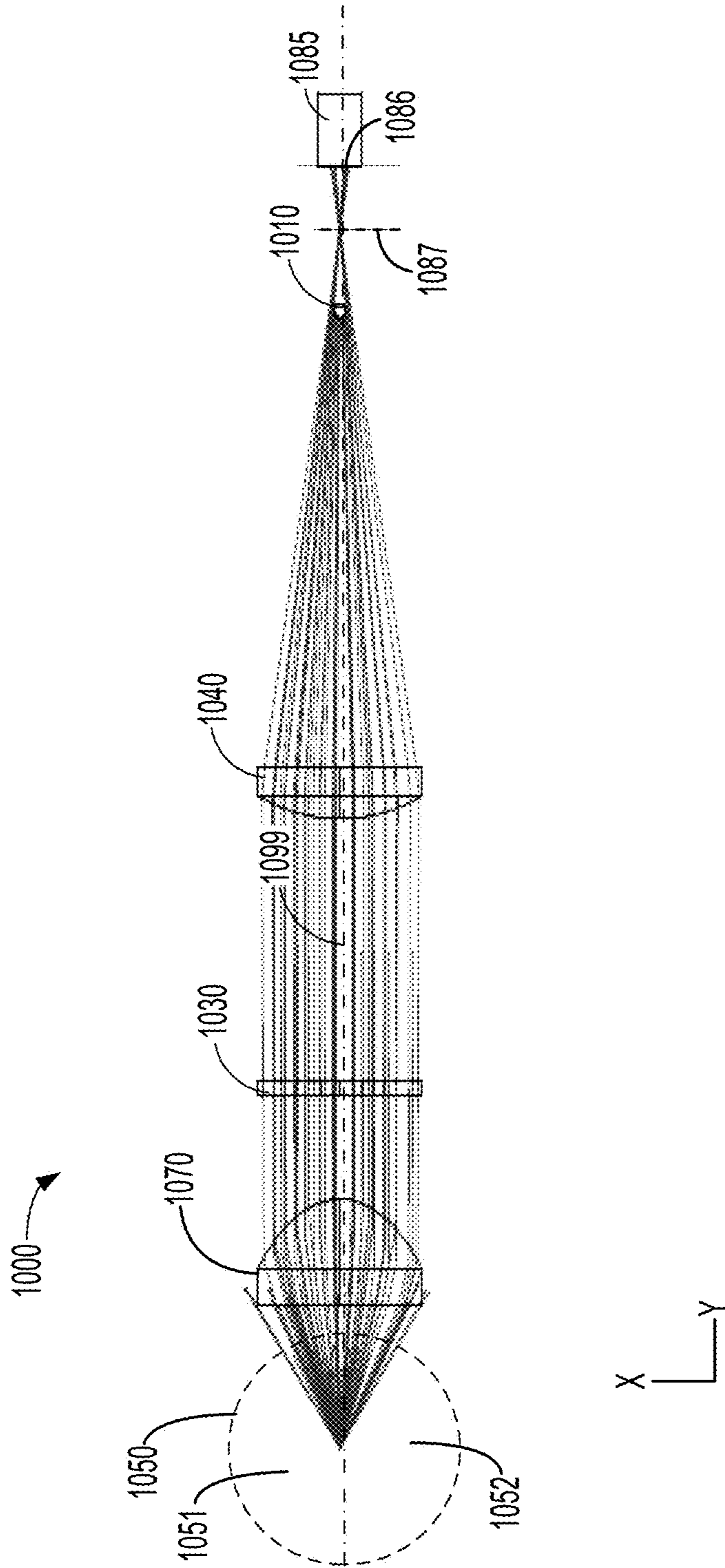


FIG. 10

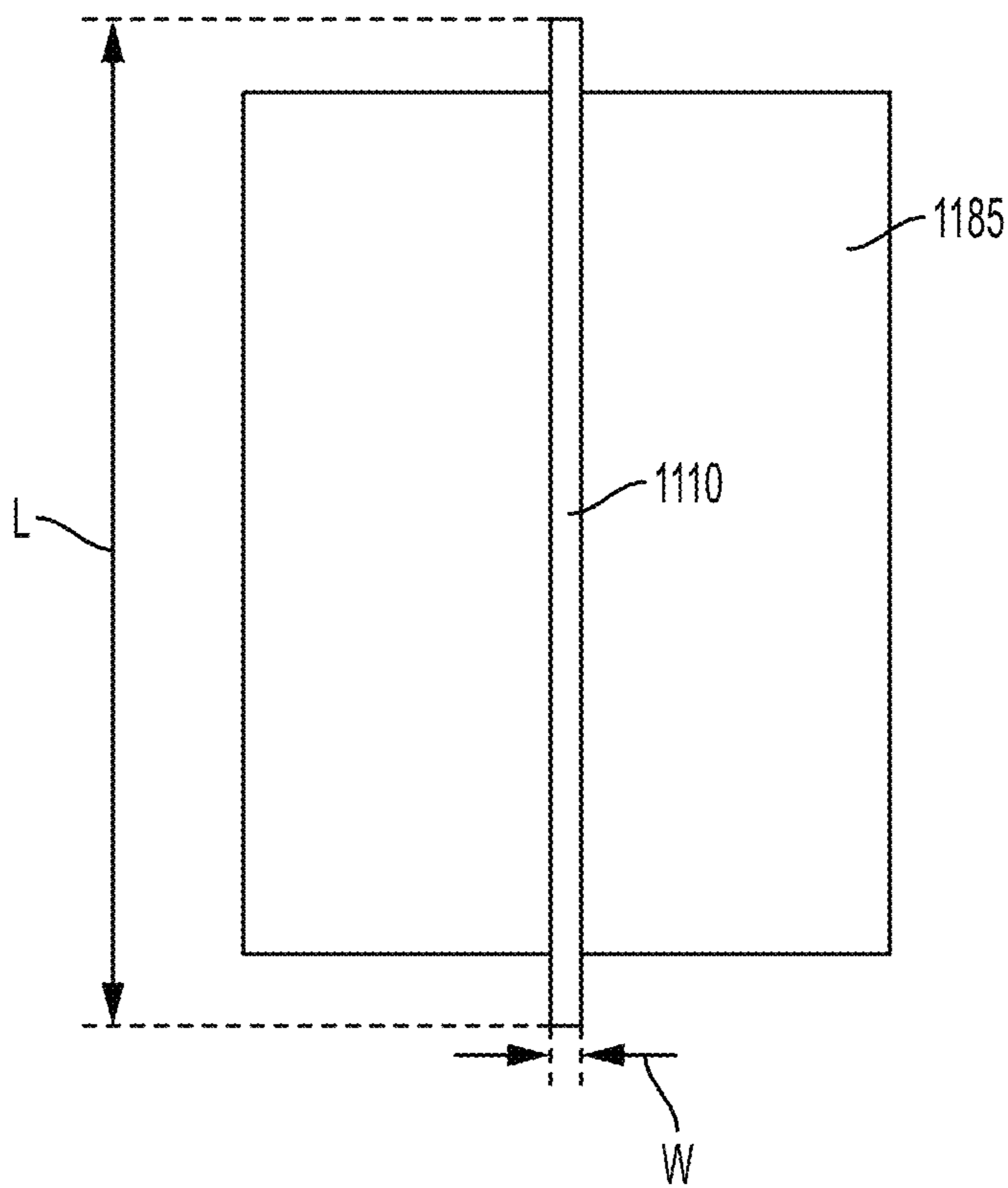


FIG. 11

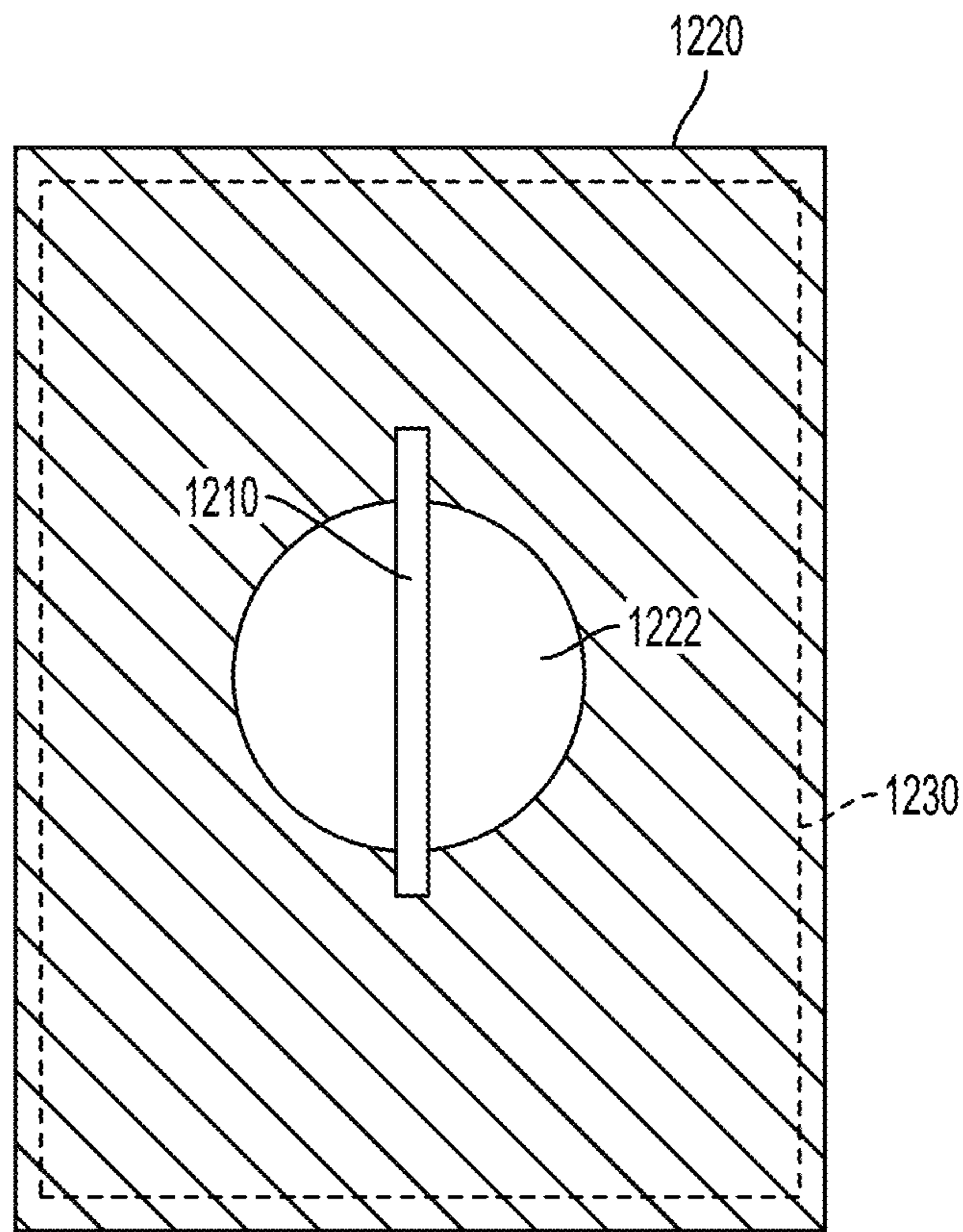


FIG. 12

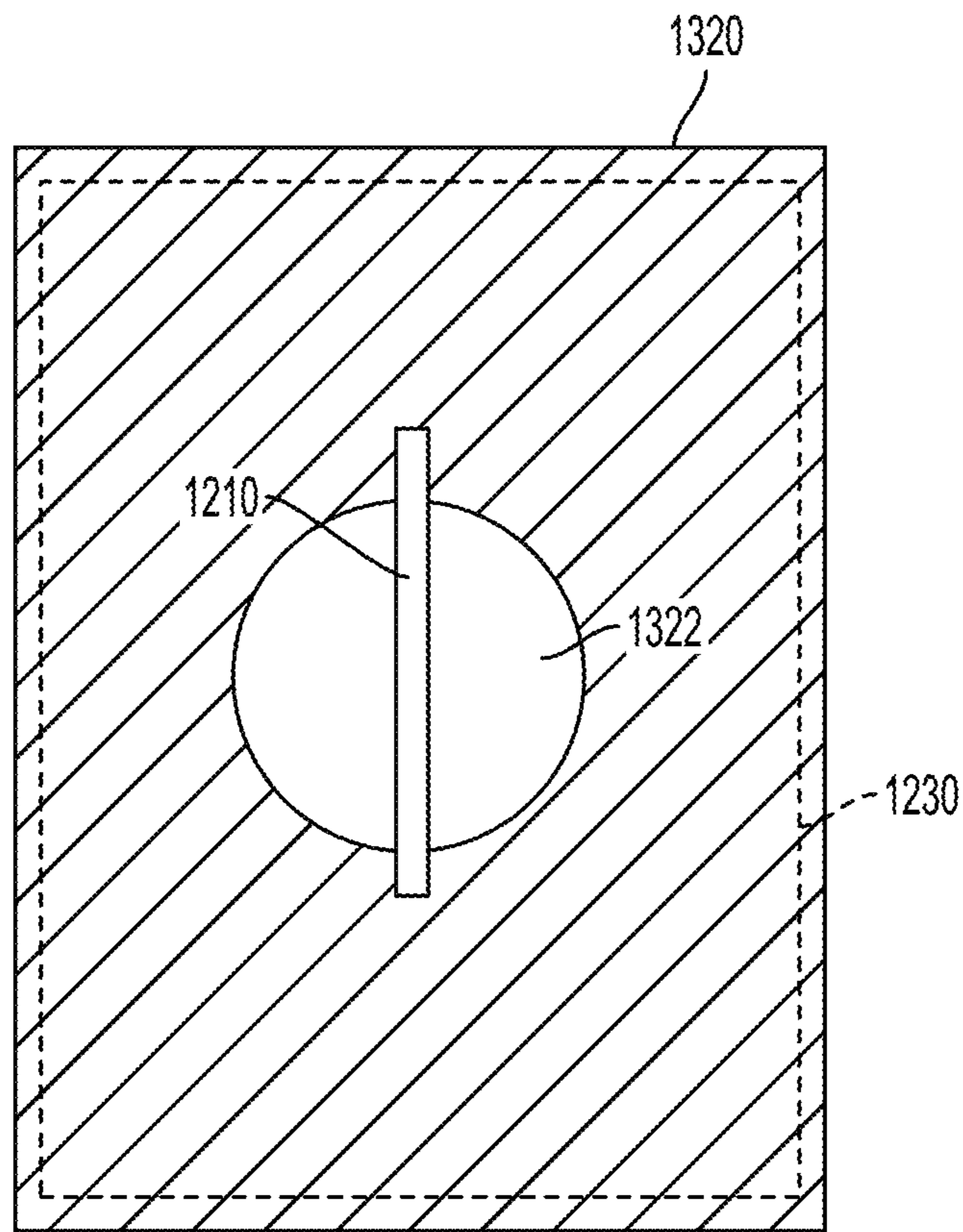


FIG. 13

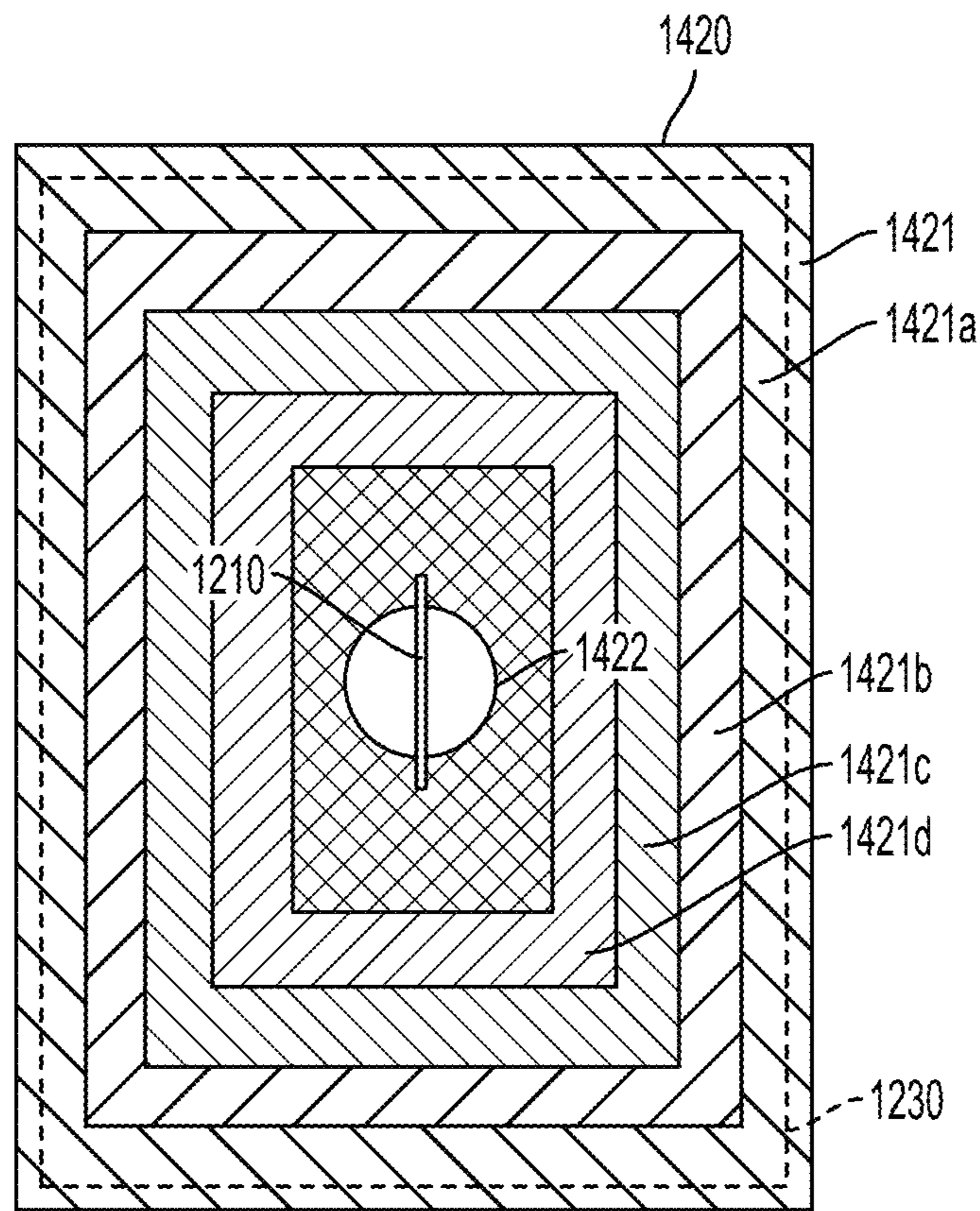


FIG. 14

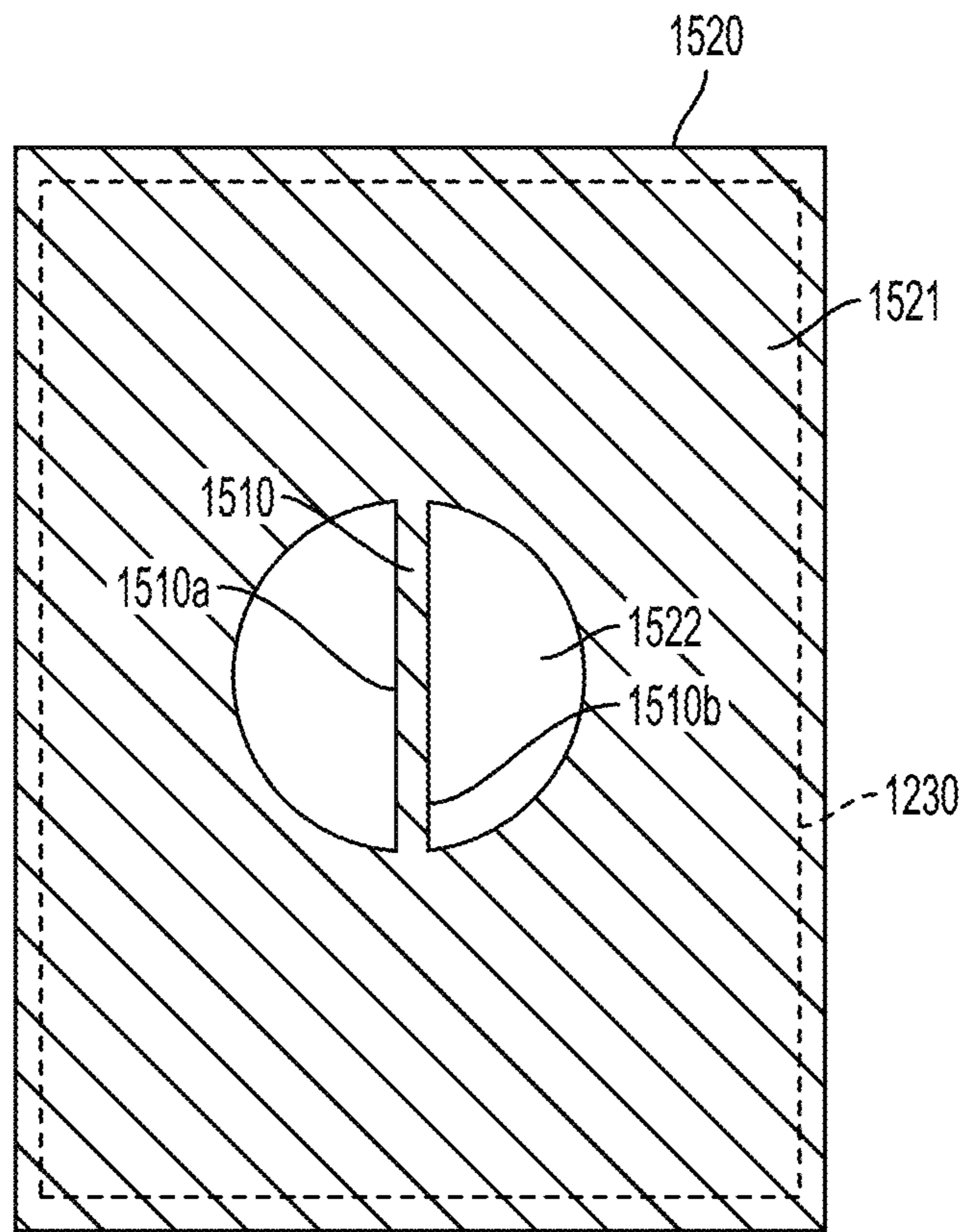


FIG. 15

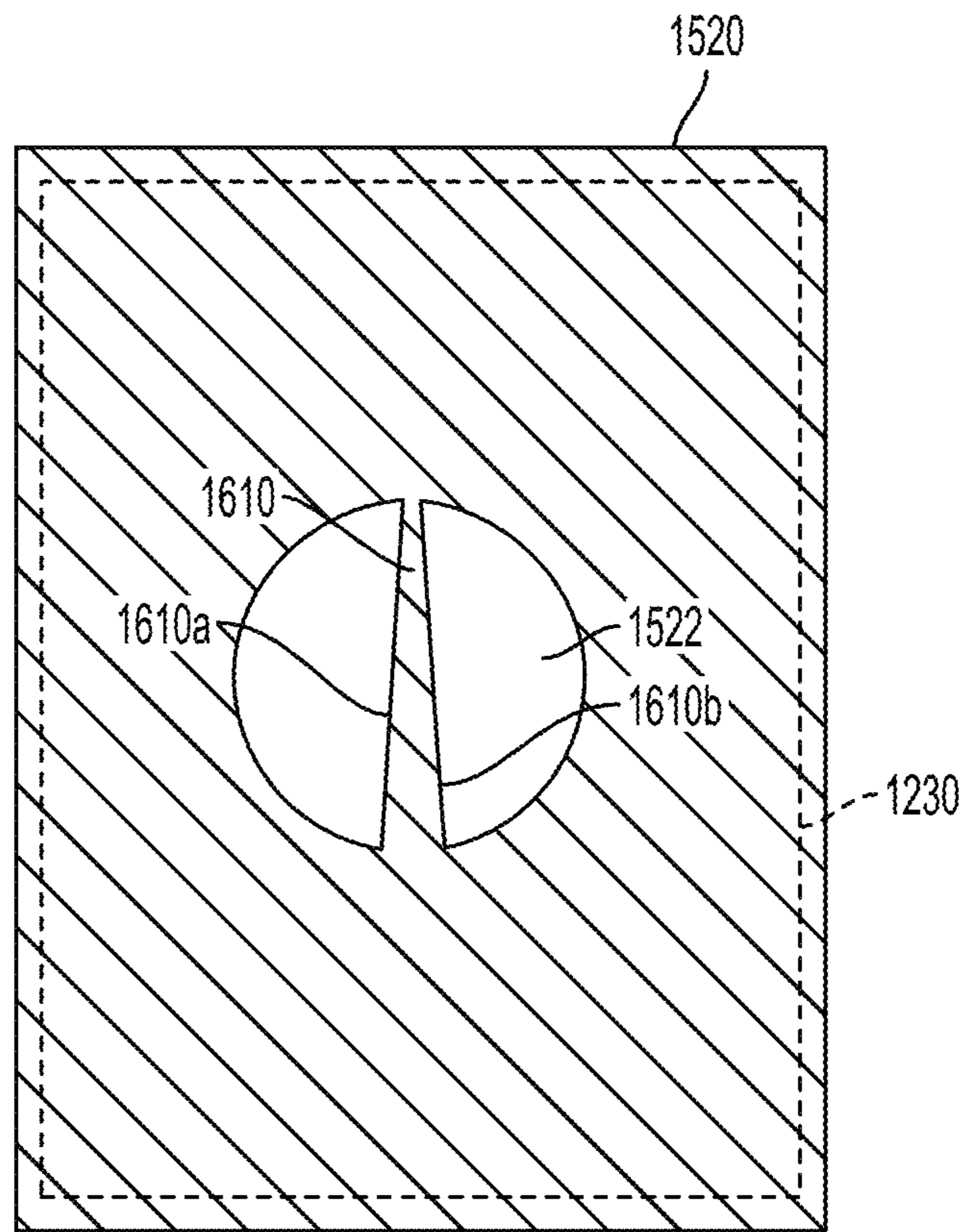


FIG. 16

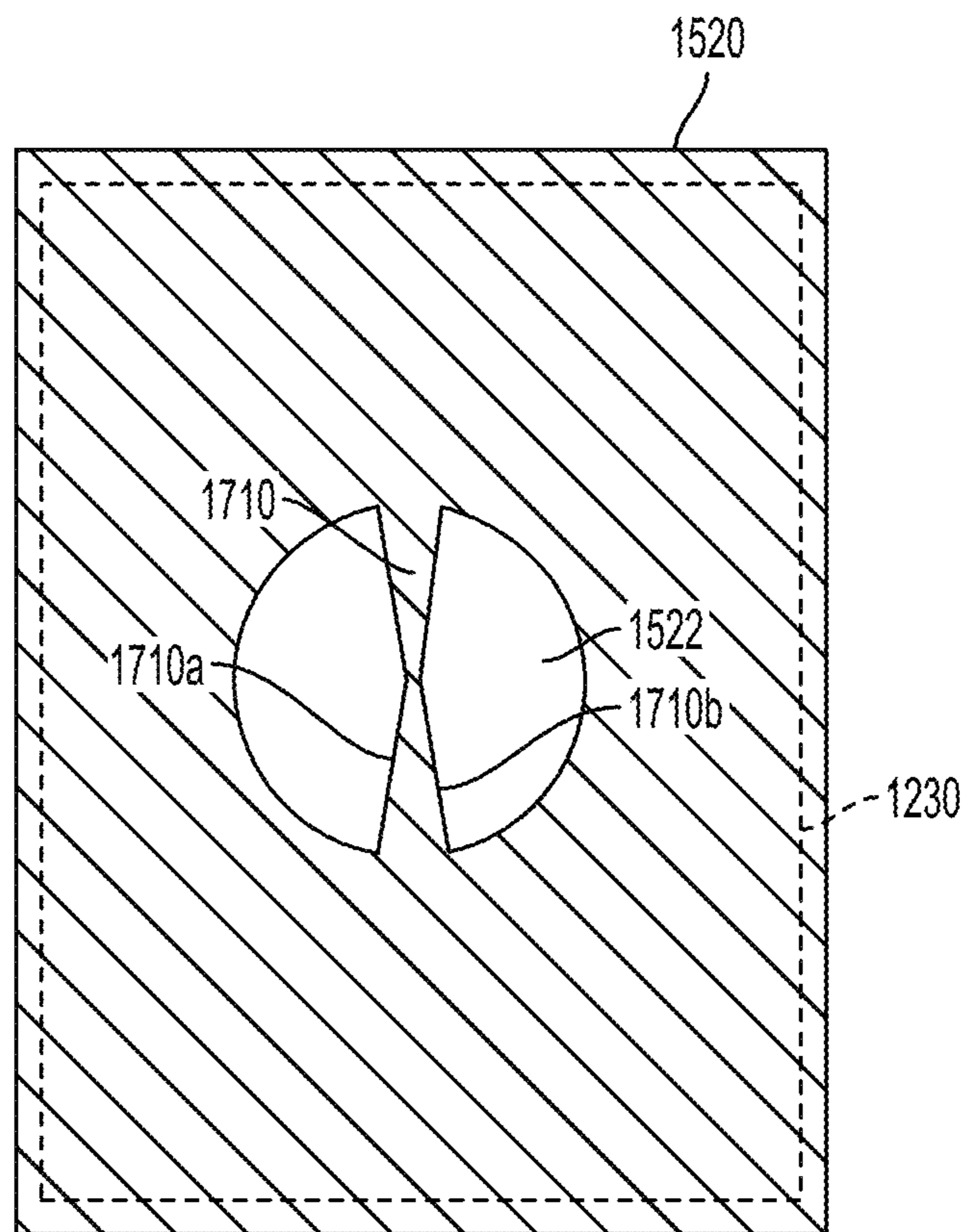


FIG. 17

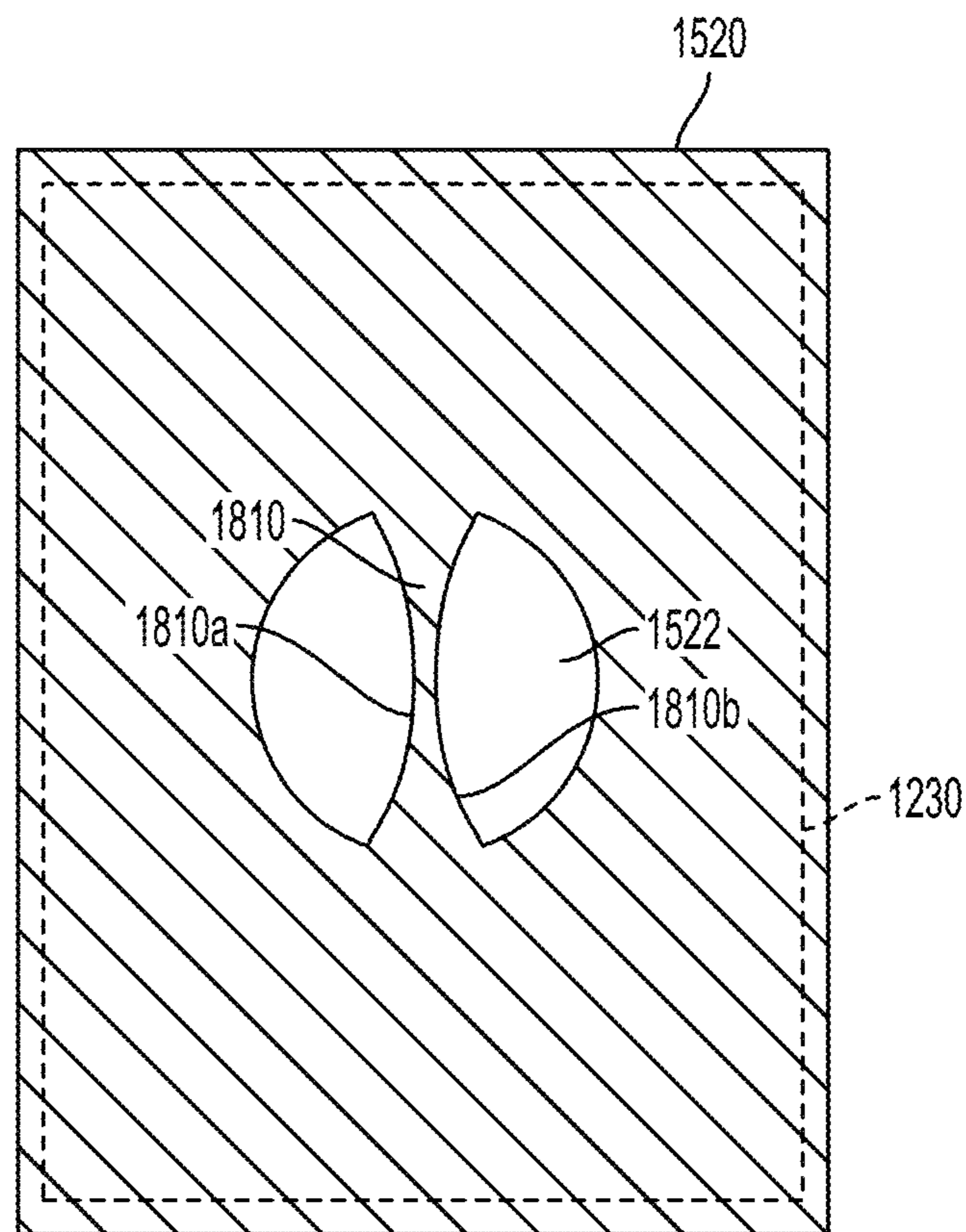


FIG. 18

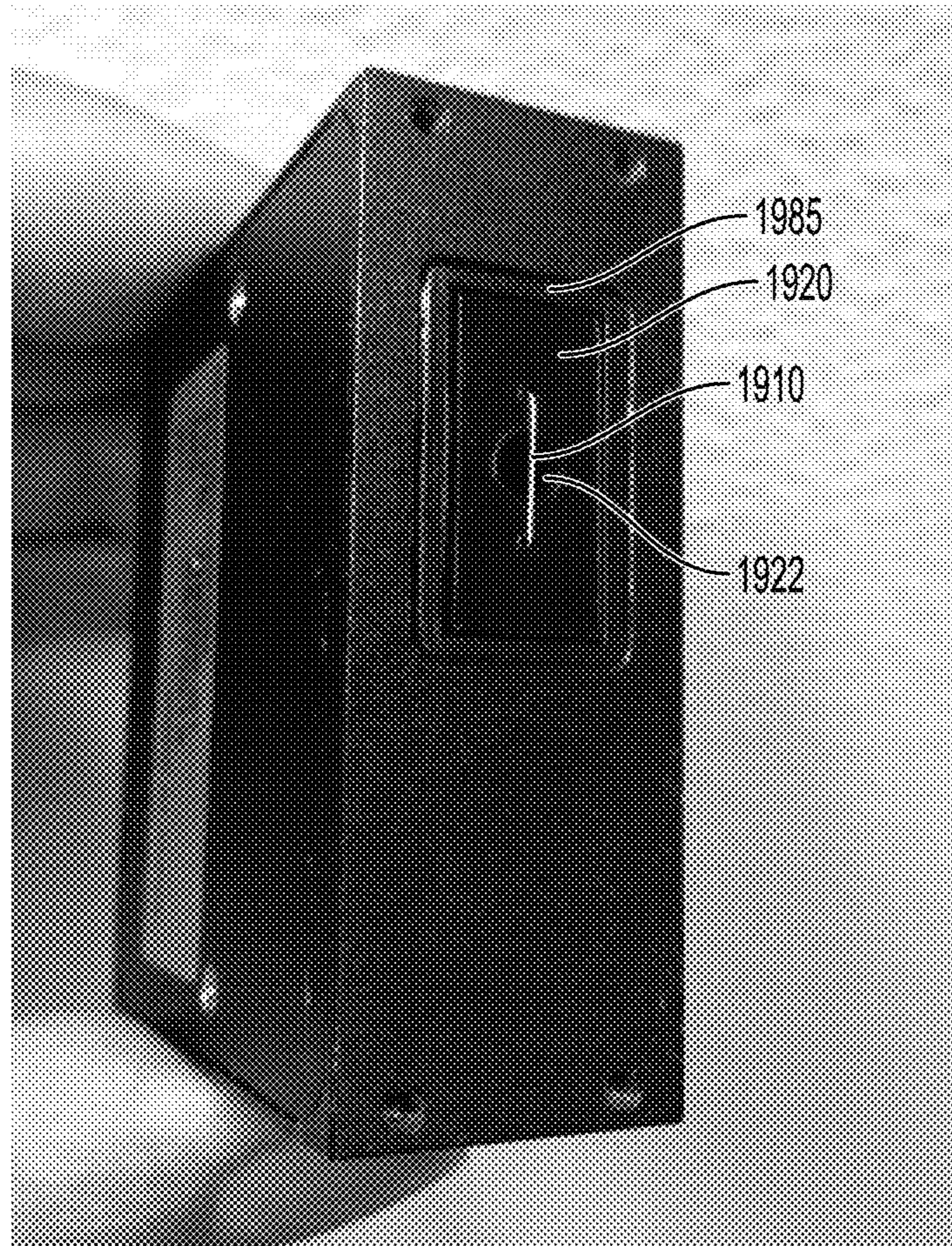


FIG. 19

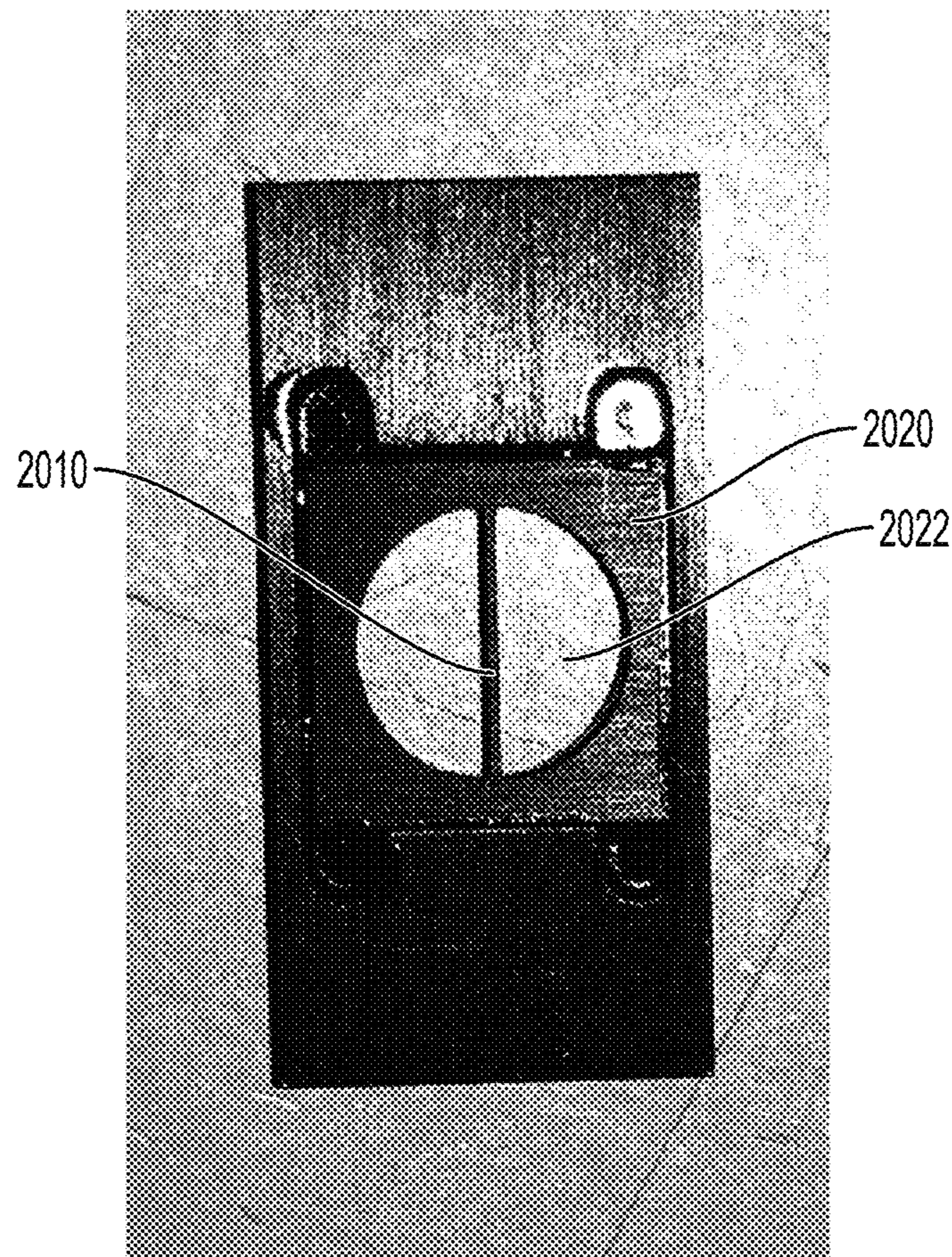


FIG. 20

LIGHT COLLECTION FROM OBJECTS WITHIN A FLUID COLUMN

BACKGROUND

Object discrimination techniques distinguish between objects of different types. These techniques are particularly useful to sort biological cells according to cell type. Some cell sorting approaches rely on light emanating from the cells to determine their type. In some implementations, cells traveling in a column of fluid are exposed to an excitation light and light emanating from the cells in response to the excitation light is detected. Cells of a first type produce output light that is different in some characteristic, e.g., wavelength and/or intensity, from cells of a second type. The differences in output light emanating from the cells can be the basis for cell type discrimination and sorting.

SUMMARY

Some embodiments are directed to an optical arrangement configured to receive output light emanating from an object disposed within a fluid column. The output light crosses an optical refraction boundary of the fluid column between the object and the optical arrangement. The optical arrangement modifies the output light such that the modified output light has an intensity that is more uniform than an intensity of the output light. For example, within a cross section of the fluid column, the intensity of the modified output light can be substantially uniform irrespective of a position of the object.

According to some embodiments, an optical apparatus includes the optical arrangement and further includes a detector that detects the modified output light and provides an electrical signal responsive to the modified output light.

In accordance with some embodiments, a discrimination system includes an excitation light source configured to generate excitation light and to direct the excitation light toward an object in a fluid column. The object emanates output light in response to the excitation light. The system comprises an optical arrangement configured to receive the output light. The output light crosses an optical refraction boundary of the fluid column between the object and the optical arrangement. The optical arrangement modifies the output light such that the modified output light has an intensity that is more uniform than an intensity of the output light, e.g., the intensity of the modified output light is substantially uniform irrespective of a position of the object in a cross section of the fluid column. An optical detector is configured to detect the modified output light and to provide an electrical signal responsive to the modified output light. Object type discrimination circuitry discriminates between a first type of object and a second type of object based on the electrical signal.

BRIEF DESCRIPTION OF THE DRAWINGS

FIG. 1A is a diagram of a system incorporating an optical apparatus in accordance with some embodiments;

FIG. 1B shows an xy plane cross section of the fluid column in the measurement region of the system of FIG. 1A;

FIG. 2A shows light emanating from an object located near the center of the fluid column with substantially no refraction of light at the fluid-air interface of the fluid column.

FIG. 2B shows light emanating from an object located in an upper portion of the elliptical core of the fluid column exhibiting refraction of light at the fluid-air interface;

FIG. 3 illustrates development of an analytical formula for angular dependence of the in-plane light ray density as a function of position x ;

FIG. 4A provides a family of graphs showing the angular dependence of radiance for different object positions;

FIG. 4B provides a family of graphs of the relative intensity of light collected from the fluid column with respect to object position along the x axis for different numerical apertures of the collection optics.

FIG. 5A provides a family of graphs of the angular dependence of radiance for different positions of the object and showing regions of exclusion;

FIG. 5B shows the relative intensity of light collected from the fluid column with respect to object position along the x axis when no angles are excluded, when rays having angles between -0.3 rad and $+0.3$ rad are excluded, and when rays having angles between -0.4 rad and $+0.4$ rad are excluded;

FIG. 6 is a flow diagram of an approach for identifying objects traveling in a fluid column with reduced positional variation of detected output light in accordance with some embodiments;

FIG. 7 is a top view of a ray tracing simulation of an optical system that includes an optical apparatus in accordance with some embodiments;

FIG. 8 is a photograph of a split objective lens configured to reduce the positional variation of detected output light for objects in a fluid column in accordance with some embodiments;

FIGS. 9A and 9B illustrate the simulated performance of the optical apparatus of FIG. 7;

FIG. 10 is a top view of a ray tracing simulation of an optical system that includes an optical apparatus in accordance with some embodiments;

FIG. 11 illustrates an optical apparatus comprising an elongated mask feature in accordance with some embodiments;

FIG. 12 depicts an optical apparatus comprising an elongated mask feature that can be used in conjunction with a plate having an aperture according to some embodiments;

FIG. 13 illustrates an optical apparatus comprising a plate and elongated mask feature wherein the optical transparency of the plate varies smoothly with position in accordance with some embodiments;

FIG. 14 illustrates an optical apparatus comprising a plate and elongated mask feature wherein the optical transparency of the plate varies with position in accordance with some embodiments;

FIG. 15 is a diagram of an optical apparatus comprising plate and an elongated mask feature extending across an aperture formed as a unitary structure in accordance with some embodiments;

FIGS. 16 through 18 illustrate various configurations of longitudinal edges of elongated mask features in accordance with several embodiments;

FIG. 19 is a photograph of an optical apparatus that includes a wire mask in accordance with some embodiments; and

FIG. 20 is a photograph of an optical apparatus that includes a plate having an elongated mask feature extending across an aperture formed as a unitary structure in accordance with some embodiments.

The figures are not necessarily to scale. Like numbers used in the figures refer to like components. However, it will be understood that the use of a number to refer to a component in a given figure is not intended to limit the component in another figure labeled with the same number.

Embodiments described herein relate to devices, systems and methods for discriminating between different types of objects. The objects emanate output light in response to an excitation light that is directed toward the objects in a fluid column, such as a flow stream. In some implementations, cell types are distinguished based on the intensity of the output light emanating from the objects. Specific embodiments discussed herein are directed to distinguishing between X chromosome sperm cells and Y chromosome sperm cells. It will be appreciated that the approaches of this disclosure can be applied more generally to distinguishing between any objects of different types so long as the output light emanating from one object type has a discernable difference in at least one characteristic when compared to the light emanating from another object type. In some examples provided, the fluid column is a flow stream that has a curved boundary or interface where refraction of light may occur. For example, the curved boundary of the fluid column may be generally circular in cross section. The fluid column can be bounded by solid walls or may be jetted into the air. The objects may move along the fluid column which may include a central core shaped by a sheath fluid that at least partially surrounds the central core. Light emanating from the objects encounters at least one optical refraction boundary between the objects and other materials, such as at the interface between the fluid column and air.

Due at least in part to the refraction at the fluid-air interface, the light collection efficiency external to the fluid column of light emanating from objects within the column depends upon the position of the objects for systems in the prior art. Light collection efficiency that varies with position is detrimental in applications where the light emanating from the objects must be precisely quantified and such precision is limited by random (not directly observable) position fluctuations of the objects. The approaches disclosed herein enhance the precision of systems that may be limited by such fluctuations, such as jet-in-air flow cytometers. As discussed in more detail below, the positional variability of light intensity collected from objects in a fluid column can be addressed by selectively masking rays at one or more planes (e.g., aperture stop, field stop) of the optical system in order to reduce the dependence of intensity on position.

The approaches outlined herein are particularly applicable to flow cytometry. However, the approaches can be applied to any system where light is collected on one side of an interface from objects emanating the light from the other side of the interface, wherein the interface causes a variation in the emanating light ray paths in a manner dependent on the object's position relative to the detector. Approaches herein modify the light collection efficiency of the output light emanating from the object to compensate for positional variation within the fluid column.

The "jet-in-air" flow cytometer system **100** illustrated schematically in FIG. 1A is one type of flow cytometer that can be used to discuss the concepts of the disclosure. The "jet-in-air" flow cytometer system **100** pumps fluid into a chamber **110** at high pressures causing a flow stream **150** comprising a fluid column to jet out of the exit nozzle **160** of the chamber **110** at high velocity, e.g., about 20 m/s. The fluid column **150** expelled from the exit nozzle **160** can be roughly circular in cross-section and may have a diameter of about 10 μm to about 100 μm in some implementations. The flow stream **150** is composed of a core stream **151** within a

sheath stream **152** where the arrows in FIG. 1A indicate the direction of flow of the core and sheath streams **151**, **152**.

Within the chamber **110**, a sample output nozzle **111** ejects the core stream **151** containing objects **171**, **172** which may be of multiple types. The core stream **151** is bounded and shaped by a stream **152** of sheath fluid which is ejected from a sheath fluid nozzle (not shown) into the chamber **110**. The sheath stream **152** at least partially surrounds the core stream **151**, and the sheath stream **152** and the core stream **151** do not substantially mix. The sloping or angled walls **115** of the chamber **110** cause the sheath stream **152** to narrow and/or maintain the cross-sectional size of the core stream **151** within the flow stream **150** before and after the flow stream **150** is ejected from the exit nozzle **160** of the chamber **110**. The movement of the sheath stream **152** constrains the objects **171**, **172** in the core stream **151** to move toward the center of the flow stream **150** when the fluid column **150** is ejected from the chamber **110**. The flow stream **150** delivers the objects **171**, **172** to a measurement region **175** of the flow stream **150**, e.g., in single file.

As the objects pass through the measurement region **175** of the flow stream **150**, light from an excitation light source **180** provides excitation light to the objects **171**, **172**. The excitation light source **180** can provide light in a broad wavelength band or in a narrow wavelength band. For example, the excitation light source **180** may be a laser. In some configurations, the excitation light may be modified by an optical element **181**. For example, the excitation light may be focused on the measurement region **175** by a lens **181**. Objects in the measurement region **175** emanate light, e.g., scattered or fluorescent light, in response to the excitation light source **180**.

Objects of a first type **171** will emanate light that differs in at least one characteristic in comparison to light that emanates from objects of the second type **172**. For example, in some scenarios, objects of the first type **171** will emanate light having a higher intensity than the light that emanates from objects of the second type **172**.

An optical collection arrangement **190** is arranged to collect the output light **161** emanating from the object within the measurement region **175** that crosses the optical refraction boundary of the flow stream **150** at the fluid-air interface. The optical arrangement **190** is configured to modify the output light **161** to provide modified output light **162** that compensates for position dependence of the light emanating from the object **172a** in the measurement region **175** as discussed in more detail below. A detector **185** receives the modified output light **162** and, in response, generates an electrical signal. In some scenarios, the amplitude of the electrical signal may be different for different object types. The electrical signal is used by discrimination circuitry **187** to distinguish between different types of objects **171**, **172**. For example, the discrimination circuitry **187** may be configured to compare the amplitude of the electrical signal to a threshold to discriminate between objects of the first type **171** and objects of the second type **172**.

FIG. 1B shows an xy plane cross section of the flow stream **150** in the measurement region **175**. In the xy cross section of the measurement region **175**, the core stream **151** is elliptical in shape, and the fluid of the core stream **151** comprises at least one object **172a** suspended in a buffer solution. The sheath stream **152** is substantially surrounds the core stream **151**. In a particular example used for this discussion in this disclosure, the objects **171**, **172** are sperm cells and the system **100** is implemented to discriminate X chromosome sperm from Y chromosome sperm.

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A focused laser beam generated by the excitation source **180** illuminates the sperm cell **172a** within the measurement region **175**. The cells **171**, **172** are stained with a fluorescent dye, and the excitation light causes the cell **172a** within the measurement region to emanate fluorescent output light. The purpose of the elliptical core **151** is to orient a sperm cell **172a** such that the flat sides of the sperm cell are facing to the left and the right as shown in FIG. **1B**. In this orientation, the flat sides of the sperm cell **172a** face the laser **180** and the optical collection arrangement **190**, respectively.

When the core stream **151** is elliptical, a sperm cell **172a** can take any number of positions along the x-axis within the core stream **151**. FIG. **1B** shows three possible positions for the sperm cell **172a** in the elliptical core **151**. In the orientation shown in FIG. **1B**, the first possible position for the sperm cell **172a** in the core stream **151** is approximately at the center of the elliptical core **151** (on the optical axis **199** of the optical collection arrangement **190**), a second possible position is at the top of the core stream **151** (above the optical axis **199**), and a third possible position is at the bottom of the core stream **151** (below the optical axis **199**). A position-dependent refraction of the output light rays emanating from the sperm cell **172a** occurs at the fluid-air interface **153** at the different positions within the core stream **151**.

When the sperm cell **172a** is located at the first position and the flow stream **150** has a circular cross section as shown in FIG. **1B**, the in-plane rays of light emanating from the sperm cell **172a** are approximately normally incident on the fluid-air interface **153**. Rays that emanate from points of the sperm cell **172a** away from its center, or rays that emanate out of the plane of the figure, are not exactly normally incident on the interface **153**; these rays are not considered in this simplified discussion, but one of ordinary skill in the art can see how the discussion could be generalized to include them. Thus, no refraction of light occurs at the fluid-air interface **153**.

The diagram of FIG. **2A** shows the absence of light refraction of the output light **298** emanating from a sperm cell **172a** and crossing the interface **153** when the sperm cell **172a** is at the 1st position within the elliptical core **151** shown in FIG. **1B**. Correspondingly, the in-plane density of the light rays **298** exiting the flow stream **150** in FIG. **2A** is uniform with respect to ray angle. Uniform angular density of light rays corresponds to uniform radiance as a function of ray angle.

In contrast, when a sperm cell **172a** is off the optical axis **199** and is nearer to the top or bottom of the elliptical core **151**, e.g., at the 2nd and 3rd positions of the elliptical core **151** shown in FIG. **1B**, at least some of the output rays emanating from the sperm cell **172a** encounter the fluid-air interface **153** at an oblique angle. These output rays are refracted at the fluid-air interface **153** in contrast to the normal incidence scenario discussed above. The most oblique rays are the most severely refracted. Refraction of the light rays causes the radiance distribution of the fluorescent light exiting the flow stream **150** across the fluid-air interface **153** to become non-uniform and to vary with position of the cell **172a** along the x axis. That is, this refraction changes the radiance distribution of output light emanating from sperm cell **172a** outside of the flow stream **150**.

For example, when the cell **172a** is located off the optical axis **199**, e.g., at the 2nd or 3rd positions shown in FIG. **1B**, the density of light rays and thus the radiance on the air side of the interface **153** is higher at positive or negative ray angles, respectively, with respect to the optical axis **199** when compared to the radiance on the air side of the

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interface **153** at angles parallel to the optical axis **199** or at negative or positive ray angles, respectively. Positive and negative refer to the sign of the ray angle γ in FIG. **3**. FIG. **2B** is a diagram illustrating light rays **299** emanating from a cell **172a** and exiting the flow stream **150** through the fluid-air interface **153** when the cell **172a** is located at the 2nd position of the elliptical core **151**. In this scenario, the density of light rays, or radiance, at positive ray angles is greater than the density of the light rays parallel to optical axis **199** or at negative ray angles. For an optical system with a predetermined numerical aperture (NA), the amount of light collected by the system from cells of the same type (e.g., the collection efficiency) may vary depending on whether the cell is in the first position or the second position. The positional dependence of the system collection efficiency leads to inaccuracies in determining cell type.

With reference to FIG. **3**, an analytical formula for the light ray density as a function of ray angle γ and sperm position x is determined using Snell's law, where γ is the angle of a light ray, with respect to the optical axis, emanating from the object after refraction at the fluid-air interface. This analysis considers only rays within, or tangential to, the two-dimensional cross-section of the flow stream.

We wish to solve for the density of the light rays with respect to the angle γ , which we can use to determine the density of rays at the entrance pupil of an optical collection system for each sperm position x . This can be written:

$$I_{\gamma}(\gamma). \quad (1)$$

For our purposes we can assume that the sperm cell emanates light uniformly in all directions, so the density of emanated light rays with respect to the angle θ is:

$$I_{\theta}(\theta)=1/\pi, \quad (2)$$

that is, uniformly distributed from

$$\theta = -\frac{\pi}{2} \text{ to } \theta = \frac{\pi}{2}.$$

By geometrical analysis:

$$\theta = \tan^{-1}\left(\frac{\cos\phi - x}{\sin\phi}\right); \quad (3)$$

$$\alpha = \pi/2 - \phi - \theta; \text{ and} \quad (4)$$

$$\gamma = \pi/2 - \phi - \beta, \quad (5)$$

wherein the angles γ , θ , ϕ , α , β , and the distance x are shown in FIG. **3**. As the flow stream has index of refraction n , Snell's law yields another relation between the angles:

$$\sin \beta = n \sin \alpha. \quad (6)$$

The density of light rays external to the interface $I_{\beta}(\beta)$ is related to the density of light rays internal to the interface $I_{\alpha}(\alpha)$ by the following formula, with $T(\alpha)$ representing the average, across both polarizations, of the transmission through the interface:

$$I_{\beta}(\beta) = I_{\alpha}(\alpha)T(\alpha)\left|\frac{d\alpha}{d\beta}\right| \quad (7)$$

The transmission is related to the Fresnel reflection coefficients for s- and p-polarization, $R_s(\alpha)$ and $R_p(\alpha)$, with the following formulas:

$$T(\alpha) = 1 - R(\alpha), \quad (8)$$

$$R(\alpha) = \frac{R_s(\alpha) + R_p(\alpha)}{2}, \quad (9)$$

$$R_s(\alpha) = \left| \frac{\cos \alpha - n \cos \beta}{\cos \alpha + n \cos \beta} \right|^2, \text{ and} \quad (10)$$

$$R_p(\alpha) = \left| \frac{\cos \beta - n \cos \alpha}{\cos \beta + n \cos \alpha} \right|^2. \quad (11)$$

Using Eq. (7) with the above and the following additional relations:

$$I_\phi(\Phi) = I_\theta(\theta) \left| \frac{d\theta}{d\Phi} \right|, \quad (12)$$

$$I_\alpha(\alpha) = I_\phi(\Phi) \left| \frac{d\Phi}{d\alpha} \right|, \text{ and} \quad (13)$$

$$I_\gamma(\gamma) = I_\beta(\beta) \left| \frac{d\beta}{d\gamma} \right|, \quad (14)$$

we have an expression for the density of rays with respect to γ :

$$I_\gamma(\gamma) = I_\theta(\theta) \left| \frac{d\theta}{d\gamma} \right| T(\gamma) \quad (15)$$

Now, the optical collection arrangement's NA is given by the sine of the maximum ray angle γ_0 , so we can solve for this angle in terms of NA:

$$\gamma_0 = \sin^{-1}(NA) \quad (16)$$

Finally, the relative collected light intensity, as a function of sperm position x , is given by integrating Eq. (15) from $-\gamma_0$ to γ_0 and normalizing by that integral value at $x=0$:

$$\text{Relative Intensity} = \frac{\int_{-\gamma_0}^{\gamma_0} I_\gamma d\gamma}{\int_{-\gamma_0}^{\gamma_0} I_\gamma(x=0) d\gamma}. \quad (17)$$

Using the formula for ray density distribution of Eq. (15), the angular dependence of ray density (radiance) for different sperm positions can be plotted as in FIG. 4A. In FIG. 4A, each of the lines represents the density of rays as a function of angle γ for a given sperm position x , where the angle γ is in radians. The plots correspond to a series of positions that lie in a range symmetric about $x=0$, (corresponding to graph 404 in FIG. 4A), which is where the ray density (radiance) is uniform as a function of angle. When x is positive (e.g., 2nd position in FIG. 1B, corresponding to graph 402), relative radiance is higher for positive ray angles γ and lower for negative ray angles γ , and the opposite is true when x is negative (e.g., 3rd position in FIG. 1B, corresponding to graph 403).

If the numerical aperture of the collection optics (optical collection arrangement 190 in FIGS. 1A and 1B) is large, e.g., approaching one, the variation in collected optical

intensity with respect to position for light emanating from an object within the elliptical core is relatively small. This is because essentially all light emanating from the object and directed to the right would be collected by the collection optics, regardless of the exact ray direction, and the total amount of emanating light is invariant to object position (given uniform excitation). In contrast, a small numerical aperture results in a relatively large collected intensity variation with respect to object position, because changes in object position affect the radiance distribution, and a small numerical aperture implies only a portion of this changing radiance distribution is collected. Practical systems may have NAs that are significantly less than one, e.g., less than 0.5, or less than 0.3. The family of graphs provided in FIG. 4B illustrates the relative intensity of light collected from an object, as a function of object position x , through collection optics with different NAs. FIG. 4A illustrates the range of angles γ captured by the different numerical apertures of FIG. 4B.

In the family of graphs of FIG. 4B, graph 412 shows the relative intensity with respect to position along the x axis for collection optics (e.g., optical collection arrangement 190 shown in FIGS. 1A and 1B) having a numerical aperture (NA) of 0.2; graph 414 shows the relative intensity with respect to position along the x axis for collection optics having an NA of 0.4; graph 416 shows the relative intensity with respect to position along the x axis for collection optics having an NA of 0.6; graph 418 shows the relative intensity with respect to position along the x axis for collection optics having an NA of 0.8; and graph 419 shows the relative intensity with respect to position along the x axis for collection optics having an NA of 0.9. It is clear from FIGS. 4A and 4B that collection optics having smaller NAs produce a larger variation in collected light intensity with respect to object position when compared to collection optics having larger NAs. Additionally, collection optics with larger NAs collect light rays having a wider range of refraction angles than collection optics having smaller NAs, and therefore have a higher overall collection efficiency.

Various embodiments disclosed herein are directed to collection devices (e.g., arrangement 190 shown in FIGS. 1A and 1B) that reduce the variation in collected light intensity with respect to object position in a flow stream. Some embodiments discussed herein can provide modified output light that has less than about a 3%, or less than about a 2%, or even less than about a 1% measured intensity variation for a deviation in position of the object that is less than 60% of a radius of the flow stream away from a center of the flow stream along an axis perpendicular to the optical axis. Many applications are sensitive to intensity measurement errors, which may arise from a variety of sources. Due to the difficulty in reducing intensity fluctuations by precisely controlling the position of objects within the flow stream, it is useful to instead reduce the variation in collected light intensity with respect to object position by careful design of the optical collection arrangement. For applications such as X/Y sperm sorting, it is often the case that two or more cell populations are to be separated based on the difference in measured fluorescence intensity between the populations. If the random position fluctuations lead to fluctuations in collected light intensity that are greater in magnitude than the nominal difference in fluorescence intensity of the two populations, it is not possible to distinguish them with simultaneously high yield and high purity. The fluorescence intensity difference between X and Y sperm cells is typically only a few percent (e.g., ~4% for bovine sperm). Current sperm sorter systems can in theory achieve

high throughput by increasing the flow rate of the core stream, but this has the effect of increasing the width of the core stream. Consequently, there would be a large uncertainty of the sperm position within the core of the flow stream. This position uncertainty and the resultant fluctuations in collected fluorescence intensity limit the maximum throughput of current sperm sorter systems to levels which do not obscure the small fluorescence intensity difference between X and Y sperm.

In sperm sorter applications, the sperm cells may be stained with Hoechst 33342 (Ho33342), a cell-permeable dye that enters the cell nuclei and binds selectively to A-T base pairs in the minor groove of double-stranded DNA within the sperm head of live cells. Typically, a UV laser is used to excite the stained sperm cells. When excited optically (at or near 350 nm), Ho33342 stained Y-chromosome bearing (male) and X-chromosome bearing (female) sperm can be resolved by measuring a small difference in total fluorescence from each cell. The difference in total fluorescence is proportional to the amount of stain within the sperm cell, which is proportional to the chromosomal content. This difference varies between mammalian species, but in domestic animals it is on the order of 4%.

One approach for intensity-position compensation is evident in FIGS. 4A and 4B. The brackets in FIG. 4A highlight regions of integration that correspond to fluorescence collection optics with a given NA. Graphs of the collected intensity variation with respect to object position for the NAs of FIG. 4A are provided in FIG. 4B. In FIG. 4B, for a given NA, integration over the fluorescence collection region is performed such that the intensity of collected light can be plotted as a function of each sperm position. It is evident from FIG. 4B that increasing the NA of the collection optics helps to decrease the influence of object position on the fluorescence intensity gathered via the collection optics.

Embodiments described herein relate to collection optics (e.g., the optical collection arrangement 190 in FIGS. 1A and 1B) that reduce collected light intensity variation with respect to object position as described above. According to some embodiments, the collection optics operate by masking rays in “angle space”, that is, the collection optics selectively collect, attenuate, and/or block rays from different angles γ in order to achieve a desired intensity vs. position profile. In practice, an “angle space” masking function can be applied at a pupil (e.g., entrance pupil, exit pupil, or aperture stop) of an optical system, where the position of a ray intersection with the pupil plane corresponds to the angle γ . In some embodiments, the collection optical arrangement achieves a desired, e.g., flatter, intensity vs. position profile by preferentially collecting higher angle (pointing away from the optical axis) light rays over lower angle light rays.

FIGS. 5A and 5B illustrate how excluding low-angle refracted rays, at a given NA, causes the intensity-vs-position curve to flatten out. Excluding the low angle rays excludes the rays that produce the most variation in the intensity vs. position profile, whereas the angular variation of radiance at high positive angles tends to cancel the corresponding variation at high negative angles. FIG. 5A shows plots of the relative radiance vs. ray angle, γ , for different positions of the object along the x axis where the angle γ is in radians. In FIG. 5A, each graph corresponds to an object position, x, within the core of a flow stream, as indicated in FIG. 3. The brackets in FIG. 5A show the portion of the light rays that will be excluded by the collection optics for each position x, when rays having angle

magnitude less than 0.3 rad are excluded (bottom bracket in FIG. 5A) and when rays having angle magnitude less than 0.4 rad are excluded (top bracket in FIG. 5A).

FIG. 5B shows the relative collected light intensity vs. position of the object along the x axis when no angles are excluded (graph 500), when rays having angles between -0.3 rad and $+0.3$ rad are excluded (graph 503) and when rays having angles between -0.4 rad and $+0.4$ rad are excluded (graph 504). Graph 5B shows that when lower angle rays are excluded, the relative intensity vs. position graph exhibits less intensity variation with respect to position.

An approach for identifying objects traveling in a fluid column in the presence of positional variation is illustrated in the flow diagram of FIG. 6. The process includes modifying 620 output light emanating from the object passing through a cross section of a flow stream such that an intensity of the modified output light is more uniform than the intensity of the unmodified output light. In some embodiments, the modified output light is substantially uniform irrespective of a position of the object. The modified output light is detected 630 and an electrical signal is generated 640 in response to the detected modified output light. A processor or other circuitry may use the electrical signal to discriminate 650 between objects of different types. For example, the circuitry may compare the amplitude of the electrical signal (which corresponds to the intensity of the detected light) to a threshold value to discriminate between objects of a first type and objects of a second type. Optionally, in some implementations, excitation light may be generated 610 by an excitation source and directed to the cross section of the flow stream wherein the object emanates the output light in response to the excitation light.

FIG. 7 is a top view of a ray tracing simulation of an optical system 700 that includes an optical apparatus 710 in accordance with some embodiments. Apparatus 710 effectively extends the NA of the collection optics in the plane of a cross section of a fluid column, preferentially collecting higher angle rays that are more balanced in terms of position vs. intensity over lower-angle rays that tend to inject variation into the position vs. intensity profile, as explained in the discussion of FIGS. 5A and 5B. Optical collection arrangement 710 modifies light emanating from an object in a cross section of a flow stream such that an intensity of the modified output light is more uniform than the output light emanating from the object. The modified output light can be substantially uniform irrespective of a position of the object within the cross section. In this particular embodiment, the intensity of the modified output light is substantially uniform irrespective to the position of the object along an axis perpendicular to an optical axis of the collection arrangement. Other embodiments may cause an intensity of the modified output light to be substantially uniform independent of a position of the object along another axis, such as the optical axis of the collection arrangement.

The optical collection arrangement 710 preferentially collects light rays emanating from the object at higher angles with respect to the optical axis 799 of the optical arrangement over light rays emanating from the object at lower angles with respect to the optical axis 799. In some implementations, the optical collection arrangement 710 is a split objective lens. A first section 711 of the split objective lens 710 collects a first portion 751 of the higher angle light emanating from the object (object not shown in FIG. 7). A second section 712 of the split objective lens 710 collects a second portion 752 of the higher angle light emanating from the object. As shown in FIG. 7, in some embodiments, a

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mask that further prevents the collection of lower angle rays may be disposed anywhere that the lower angle rays would be blocked, e.g., near an aperture stop or pupil plane, where the light is collimated. For example, a mask **786** may be disposed between the two lenses **711**, **712** as shown in FIG. 7.

As indicated in FIG. 7, the system **700** may be implemented as a folded optical system using mirrors **721**, **722**, **723**, **724** to redirect the collected portions of light along the optical axis **799** of the system **700** and toward the detector **785**. Mirrors **721**, **722** redirect the first portion **751** of light toward and along the optical axis and mirrors **723**, **724** redirect the second portion **752** of light toward and along the optical axis **799**. As illustrated in FIG. 7, the system **700** may optionally include a filter **730** such as an optical bandpass or longpass filter configured to substantially attenuate the excitation light. The system **700** can include lens **740** configured to focus the first and second portions **751**, **752** of the light toward the detector **785**.

FIG. 8 is a photograph of a split objective lens configured to reduce the variation of intensity with respect to object position. It should be noted that the split-objective design allows the fluorescence collection optics to be placed closer to the flow stream than would otherwise be possible, due to the spatial obstruction caused by the nozzle generating the flow stream. A single lens with the same effective NA as the split objective would be too big to place its focal point immediately below the nozzle generating the flow stream. The optical detection of objects within the flow stream is optimally performed immediately after the flow stream exits the nozzle, where the stream is most stable, therefore it is important to have high NA optics that do not interfere with the nozzle.

FIG. 9A illustrates the simulated performance of the split objective lens based on the model in FIG. 7 vs. the simulated performance of the comparative arrangement shown in FIG. 10 without the use of the spatial mask **1010**. Graph **901** provides the intensity with respect to object position for a system that includes the split objective lens discussed above. Graph **902** is the intensity vs. position of the comparative arrangement and is provided in FIG. 9A for comparison. Note the higher overall collection efficiency of the split-objective arrangement in addition to the decreased effect of position variation on collected light intensity.

In order to show this comparison more clearly, in FIG. 9B, each graph **901a**, **902a** is normalized to 100%. Graph **901a** provides the collection efficiency relative to center position for the split objective lens arrangement. Graph **902a** shows the collection efficiency relative to center position for the comparative system with only a single objective lens. The split objective lens yields less than 1% deviation from the center collection efficiency at an object position of 20 μm , whereas the comparative arrangement yields over 10% deviation from the center collection efficiency at the same object position.

It is possible to decrease variation in the position vs. intensity profile using other approaches as well, all of which are considered novel aspects of the disclosure. For example, rather than a collection arrangement that selects rays in angle space (e.g., close to a pupil of the optical system), as exemplified by the split objective lens, it is possible to select rays in image or position space (e.g., close to an image plane of the optical system). One example of an optical collection arrangement that selects rays in position space is a spatial mask near an image plane of the optical system. In order to make such a mask easier to align and more robust to

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alignment, it helps to increase the optical system magnification, allowing one to use larger mask feature sizes.

FIG. 10 is a top view of a ray tracing simulation of an optical system **1000** that includes an optical collection arrangement **1010**, e.g., a spatial mask, that attenuates light rays emanating from the object near the center of a flow stream cross section while not attenuating light rays emanating from the object at the top and bottom of the flow stream cross section. The term “attenuating” as used herein encompasses partially blocking or fully blocking the light rays. For example, an attenuated light ray may have an intensity reduction of 25% or 50% or 75% or even 100% when compared to its original intensity, wherein 25%, 50%, and 75% attenuation corresponds to a light ray that is partially blocked and 100% attenuation corresponds to a light ray that is fully blocked. The system **1000** shown in FIG. 10 includes a single objective lens **1070** that collimates light emanating from the object (object not shown in FIG. 10). System **1000** optionally includes a filter **1030**, such as a bandpass filter or a longpass filter configured to block excitation light from reaching the detector **1085**. A lens **1040** can be used to focus collected light toward the sensitive region **1086** of the detector **1085**. A spatial mask **1010** attenuates or blocks light rays emanating from the center of the flow stream cross section **1050** from reaching the detector **1085** while not attenuating or blocking light rays emanating from the top and bottom regions **1051**, **1052** of the flow stream cross section **1050** from reaching the detector **1085**. In FIG. 10, the top region **1051** of the flow stream cross section **1050** refers to the portion of the flow stream cross section that is above the optical axis **1099** in FIG. 10. The bottom region **1052** of the flow stream cross section **1050** is below the optical axis **1099** in FIG. 10. The mask **1010** may be opaque to the emanating light or may be semi-transparent. In some embodiments, the optical transparency of the mask **1010** may vary with position, e.g., such that an image of the center of the flow stream cross section is more attenuated than an image of the top and/or bottom of the flow stream cross section.

A simple spatial mask that alleviates the variation in collected intensity with object position is a thin wire (for example, having a diameter in a range of about 100 microns to 300 microns, e.g., about 200 microns in diameter) in front of the optical detector **1085** and near an image of the flow stream, with the wire axis oriented parallel to the flow stream and nominally centered with respect to the optical axis **1099**. The effect of the wire can be varied by moving it in and out of the image plane **1087**, which is where an image of the flow stream appears (a magnified image in the current embodiment).

A spatial mask that reduces the variation in collected intensity with object position is illustrated in FIG. 11. The spatial mask comprises an elongated feature **1110**, which may be implemented as a thin wire having a circular cross sectional area, a bar having a rectangular cross sectional area or other mask feature disposed at least partially across the active area **1185** of the optical detector and near an image of the flow stream. The elongated mask feature **1110** can be implemented in many ways, including an extruded metal wire, an etched metal feature, human or animal hair, a trace deposited on a glass slide, or an ink line printed on a transparent medium.

Generally, the length of the mask feature, L , is much greater than its width, W . The mask feature **1110** may be oriented such that the length of the mask feature **1110** runs parallel to the flow stream and the mask feature **1110** is nominally centered with respect to the optical axis of the

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system (see FIG. 10). In some embodiments, the mask feature 1110 may have a width of about 100 microns to about 300 microns e.g., about 200 microns.

As illustrated in FIG. 12, an elongated mask feature 1210 can be used in conjunction with a plate 1220 having an aperture 1222 wherein the elongated mask feature 1210 is disposed at least partially across the aperture 1222 as illustrated in FIG. 12. The plate 1220 can be particularly useful for alignment of the system optics to achieve the optimal intensity difference between objects of two types with slightly different intensities.

In some embodiments, the plate may be made of a material that partially blocks (blocks more than 25% and less than 75% of the light), substantially blocks (blocks more than 75% of the light), or completely blocks (blocks 100% of the light) the light emanating from the object under test from reaching the active area 1230 of the detector. The aperture 1222 in the plate 1220 transmits substantially all of the light emanating from the object to the active area 1230 of the detector. The plate 1220 and aperture 1222 facilitates alignment of the system optics, allowing the operator to align the mask feature 1210 such that optimal contrast is achieved between the lower light intensity emanating from a first type of object and the slightly higher light intensity emanating from a second type of object

FIG. 13 illustrates another embodiment in which the optical transparency of the plate varies across its length and width. In this example, the plate 1320 is more optically transparent closer to the aperture 1322 and is less transmissive farther from the aperture. However, the opposite scenario is also possible, wherein the plate is less transmissive near the aperture and more transmissive farther from the aperture.

FIG. 14 illustrates another version of a plate 1420 that has a stepwise optical transparency gradient. The plate 1420 becomes more optically transmissive closer to the aperture 1422 in distinct steps in regions 1421a, 1421b, 1421c, 1421d. Comparing the plate 1320 of FIG. 13 with the plate 1420 of FIG. 14, the optical transparency of plate 1320 makes a gradual transition from the outer edges of the plate 1320 having lower optical transparency transitioning to a higher optical transparency nearer the aperture 1322 in the center of the plate 1320.

In some embodiments, the aperture plate and the elongated mask feature extending across the aperture are formed as a unitary structure as illustrated in FIG. 15. FIG. 15 shows a plate 1520 including an elongated feature 1510 that bisects the aperture 1522. In some embodiments the unitary aperture plate 1520 can have a optical transparency gradient as previously discussed and illustrated with reference to FIGS. 13 and 14. A unitary aperture plate as in FIG. 15 can be formed, for example, by photoetching a metal plate.

The longitudinal edges 1510a, 1510b of the elongated mask feature 1510 need not be parallel as they are shown in FIG. 15. In some embodiments the alignment process may be enhanced by having an elongated mask feature 1610, 1710 with non-parallel longitudinal edges 1610a, 1610b, 1710a, 1710b as shown in FIGS. 16 and 17. In some embodiments the longitudinal edges 1810a, 1810b of the elongated mask feature 1810 may be curved as illustrated in FIG. 18.

FIG. 19 is a photograph showing an optical arrangement comprising a plate 1920 having an aperture 1922 in accordance with some embodiments. An elongated mask feature 1910 comprising a thin wire is positioned across the aperture 1922 in front of the entrance to a photomultiplier tube detector. The wire modifies the output light emanating from

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objects in the flow stream by preferentially attenuating light emanating from objects at the center of the flow stream cross section as previously discussed. The preferential attenuation of light provides a more uniform light intensity vs. object position profile when compared to the unmodified output light.

FIG. 20 is photograph showing an optical arrangement in accordance with some embodiments comprising a plate 2020 and elongated mask feature 2010 extending across an aperture 2022 formed as one unitary structure. For example, the plate and elongated mask feature 2010 may be formed by photoetching the (split) aperture 2022.

The foregoing description of various embodiments has been presented for the purposes of illustration and description and not limitation. The embodiments disclosed are not intended to be exhaustive or to limit the possible implementations to the embodiments disclosed. Many modifications and variations are possible in light of the above teaching.

The invention claimed is:

1. An optical apparatus, comprising:
 - an optical arrangement configured to receive output light emanating from an object disposed within a fluid column, the output light crossing an optical refraction boundary of the fluid column between the object and the optical arrangement, the optical arrangement configured to modify the output light such that the modified output light has an intensity that is more uniform than the output light by preferentially attenuating light received by the optical arrangement from a first position in the fluid column compared to light received from a second position in the fluid column, wherein the second position is farther from the center of the fluid column than the first position along an axis perpendicular to the optical axis of the optical arrangement; and
 - a detector configured to detect the modified output light and to provide an electrical signal responsive to the modified output light.
2. The optical apparatus of claim 1, wherein the intensity of the modified output light is substantially uniform across a cross section of the fluid column.
3. The optical apparatus of claim 1, further comprising one or more of:
 - an excitation light source arranged to provide excitation light to the object within the fluid column, the object emanating the output light in response to the excitation light;
 - an optical filter configured to substantially block the excitation light from reaching the detector; and
 - a focusing lens arranged to collect the modified output light and to focus the modified output light toward the detector.
4. The optical apparatus of claim 1, further comprising electrical circuitry configured to receive the electrical signal from the detector and to distinguish between an object of a first type and an object of a second type based on the electrical signal.
5. The optical apparatus of claim 4, wherein:
 - an amplitude of the electrical signal is responsive to an intensity of the modified output light; and
 - the electrical circuitry is configured to distinguish between the first type of object and the second type of object by comparing the amplitude of the electrical signal to a threshold value.
6. An optical arrangement configured to receive output light emanating from an object disposed within a fluid column, the output light crossing an optical refraction

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boundary between the object and the optical arrangement, the optical arrangement configured to modify the output light such that the modified output light has an intensity that is substantially uniform irrespective of a position of the object,

wherein the optical arrangement is configured to preferentially attenuate light received by the optical arrangement from a first position in the fluid column compared to light received from a second position in the fluid column, wherein the second position is farther from the center of the fluid column than the first position along an axis perpendicular to the optical axis of the optical arrangement.

7. The optical arrangement of claim 6, wherein the optical arrangement comprises a mask configured to attenuate light from the first position in the fluid column.

8. The optical arrangement of claim 7, wherein the mask comprises a wire.

9. The optical arrangement of claim 6, wherein the optical arrangement is configured to preferentially collect a first portion of the output light emanating from the object at larger angles with respect to the optical axis relative to a second portion of the output light emanating from the object at smaller angles with respect to the optical axis.

10. The optical arrangement of claim 9, wherein the optical arrangement comprises a split objective lens configured to preferentially collect the first portion of the output light.

11. The optical arrangement of claim 10, wherein:
the first portion of light includes:
output light emanating from the object at positive angles with respect to the optical axis of the optical arrangement; and
output light emanating from the object at negative angles with respect to the optical axis of the optical arrangement; and

the split objective lens comprises:

a first lens configured to receive the light emanating from the object at positive angles with respect to the optical axis of the optical arrangement; and
a second lens configured to receive the light emanating from the object at negative angles with respect to the optical axis of the optical arrangement.

12. The optical arrangement of claim 8, wherein the optical arrangement further includes a mask configured to attenuate the second portion of the output light.

13. A discrimination system, comprising:

an excitation light source configured to generate excitation light and to direct the excitation light toward an object in a fluid column, the object emanating output light in response to the excitation light;

an optical arrangement configured to receive the output light, the output light crossing an optical refraction boundary of the fluid column between the object and the optical arrangement, the optical arrangement configured to modify the output light such that the modified output light has an intensity that is more uniform than the output light, wherein the optical arrangement comprises an elongated mask feature that attenuates the output light received from the first position, a length of the elongated mask feature being much greater than a width of the elongated mask feature, the elongated mask feature disposed proximate to an active area of the detector such that a longitudinal axis of the elongated mask feature is substantially parallel to a direction of flow of the objects in the fluid column;

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an optical detector configured to detect the modified output light and to provide an electrical signal responsive to the modified output light; and

object type discrimination circuitry configured to discriminate between a first type of object and a second type of object based on the electrical signal.

14. The system of claim 13, wherein the intensity of the modified output light is substantially uniform irrespective of a position of the object.

15. The system of claim 13, wherein the circuitry is configured to perform X/Y sperm cell discrimination.

16. The system of claim 13, wherein the elongated mask feature has a round cross sectional area.

17. The system of claim 13, wherein the elongated mask feature has a rectangular cross sectional area.

18. The system of claim 13, wherein the elongated mask feature is a metal wire.

19. The system of claim 13, wherein the elongated mask feature is a trace deposited on glass.

20. The system of claim 13, wherein the elongated mask feature is a filament.

21. The system of claim 13, wherein the elongated mask feature is an ink line printed on transparent paper.

22. The system of claim 13, wherein the elongated mask feature is disposed at about a center of an active area of the detector.

23. A method, comprising:

receiving, by an optical apparatus, output light from an object in a fluid column that has crossed an optical refraction boundary of the fluid column;

modifying the output light such that the modified output light has an intensity that is substantially uniform irrespective of a position of the object in the fluid column by attenuating a first portion of the output light received from a first position in the fluid column relative to a second portion of output light received from a second position in the fluid column, wherein the second position is farther from the center of the fluid column than the first position along an axis perpendicular to the optical axis of the optical apparatus;

detecting the modified output light; and

generating an electrical signal in response to the modified output light.

24. The method of claim 23, wherein modifying the output light further comprises:

preferentially collecting a first portion of the output light emanating from the object at higher angles with respect to the optical axis relative to collecting a second portion of the output light emanating from the object at lower angles with respect to the optical axis.

25. A flow cytometer system, comprising:

an excitation light source configured to generate excitation light and to direct the excitation light toward an object in a fluid column, the object emanating output light in response to the excitation light;

an optical arrangement configured to receive the output light, the output light crossing an optical refraction boundary of the fluid column between the object and the optical arrangement, the optical arrangement configured to modify the output light such that the modified output light has an intensity that is more uniform than an intensity of the output light, the optical arrangement comprising a plate that includes an aperture and an elongated mask feature disposed at least partially across the aperture, the elongated mask feature arranged such that a longitudinal axis of the elongated mask feature is substantially parallel to a direction of

flow of the objects in the fluid column, the plate and the elongated mask feature being less transmissive to the output light than the aperture;

an optical detector disposed adjacent to the plate and arranged to receive the modified output light transmitted 5 through the aperture, the optical detector configured to detect the modified output light and to provide an electrical signal responsive to the modified output light; and object type discrimination circuitry configured to discriminate between a first type of object and a second type of 10 object based on the electrical signal.

26. The system of claim **25**, wherein the plate and the elongated mask feature are one unitary structure.

27. The system of claim **25**, wherein the elongated mask feature is attached to the plate. 15

28. The system of claim **25**, wherein optical transparency of the plate to the output light varies as a function of position across the plate.

29. The system of claim **25**, wherein opposing longitudinal edges of the elongated mask feature are substantially 20 parallel to one another.

30. The system of claim **25**, wherein opposing longitudinal edges of the elongated mask feature are not parallel to one another.

31. The system of claim **25**, wherein opposing longitudinal 25 edges of the elongated mask feature are tapered along a longitudinal axis of the elongated mask feature.

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